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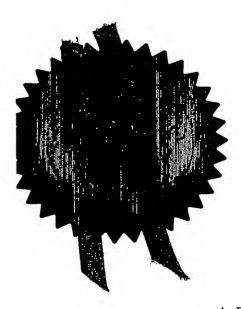
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# Binding Partners for the Thyrotropin Receptor and uses thereof

The present invention is concerned with binding partners (such as monoclonal antibodies) for the thyrotropin receptor (TSH receptor or TSHR) and uses thereof.

Thyrotropin or thyroid stimulating hormone (TSH) is a pituitary hormone which plays a key role in regulating the function of the thyroid. Its release is stimulated by the hormone TRH formed in the hypothalamus and TSH controls the formation and release of the important thyroid hormones thyroxine (T4) and tri-iodothyronine (T3). On the basis of a feedback mechanism, the thyroid hormone content of serum controls the release of TSH. The formation of T3 and T4 by thyroid cells is stimulated by TSH by a procedure in which the TSH released by the pituitary binds to the TSH receptor of the thyroid cell membrane.

In Graves' disease (a common autoimmune disorder) TSH receptor antibodies (TRAb) are formed and these autoantibodies bind to the TSH receptor in such a way as to mimic the actions of TSH, stimulating the thyroid gland to produce high levels of thyroid hormones. These autoantibodies are described as having stimulating activity. In some patients, autoantibodies bind to the TSH receptor but do not stimulate thyroid hormone production and are described as having blocking activity. (J Sanders, Y Oda, S-A Roberts, M Maruyama, J Furmaniak, B Rees Smith; "Understanding the thyrotrophin receptor function-structure relationship" Ballière's Clinical Endocrinology and Metabolism; Ed TF Davies 1997; 11(3): 451-479; pub Ballière Tindall, London).

Measurements of TSH receptor antibodies are important in the diagnosis and management of Graves' disease and other thyroid disorders. Currently three types of assay are used to measure TSH receptor antibodies:-

(a) competitive binding assays which measure the ability of TSH receptor antibodies to inhibit the binding of TSH to preparations of TSH receptor;

- (b) bioassays which measure the ability of TSH receptor antibodies to stimulate cells expressing the TSH receptor in culture; and
- (c) immunoprecipitation of TSH receptor preparations with TSH receptor antibodies.

Measurement of TSH receptor antibodies using such assays are described in references:-

J Sanders, Y Oda, S-A Roberts, M Maruyama, J Furmaniak, B Rees Smith; "Understanding the thyrotrophin receptor function-structure relationship" Ballière's Clinical Endocrinology and Metabolism; Ed TF Davies 1997; 11(3): 451-479; pub Ballière Tindall, London.

J Sanders, Y Oda, S Roberts, A Kiddie, T Richards, J Bolton, V McGrath, S Walters, D Jaskólski, J Furmaniak, B Rees Smith; "The interaction of TSH receptor autoantibodies with <sup>125</sup>I-labelled TSH receptor"; Journal of Clinical Endocrinology and Metabolism 1999; **84**(10): 3797-3802.

It has been recognised for many years that human monoclonal antibodies to the TSH receptor derived from patients' lymphocytes would be valuable reagents for understanding the pathogenesis of Graves' disease and for developing new methods of measuring TSH receptor antibodies for example as replacements for TSH in competitive binding assays. Also, as the patient's serum TSH receptor antibodies are usually powerful thyroid stimulators (TSH agonists) stimulating human monoclonal TSH receptor antibodies would be valuable for in vivo applications when tissue containing the TSH receptor (eg thyroid tissue or thyroid cancer tissue) required stimulation. Furthermore, as some patient serum TSH receptor antibodies are powerful TSH antagonists (blocking antibodies) human monoclonal TSH receptor antibodies which are TSH antagonists would be valuable for in vivo applications when the activity of tissue containing the TSH receptor (eg thyroid tissue or thyroid cancer tissue) required inactivation or to be made unresponsive to TSH, TSH receptor antibodies or other stimulators.

It has also been recognised that one of the major advantages of human monoclonal TSH receptor antibodies over TSH in such in vitro and / or in vivo applications would be the relative ease with which antibodies can be manipulated. For example, manipulation of the TSH receptor binding region of the monoclonal antibodies so as to change their characteristics, such as affinity and biological characteristics including their degree of TSH agonist or antagonist activities. Also, monoclonal antibodies will have a much longer half life than TSH in vivo and this may have considerable advantages in certain in vivo applications. Furthermore, the half life of antibodies can be manipulated easily, for example antibody Fab fragments have a much shorter half life than intact IgG. These general properties of TSH receptor antibodies are described in the publications such as B Rees Smith, SM McLachlan, J Furmaniak; "Autoantibodies to the thyrotropin receptor"; Endocrine Reviews 1988; 9: 106-121; B Rees Smith, KJ Dorrington, DS Munro; "The thyroid stimulating properties of longacting thyroid stimulator yG-globulin subunits"; Biochimica et Biophysica Acta 1969; 192: 277-285; KJ Dorrington, DS Munro; "The long acting thyroid stimulator"; Clinical Pharmacology and Therapeutics 1966; 7: 788-806.

A still further advantage of human monoclonal TSH receptor antibodies could be in their use to identify and provide new types of TSH receptor antibody binding sites. For example by the generation of antibodies to the regions of the human monoclonal TSH receptor antibodies which bind the TSH receptor. Some of the anti-idiotypic antibodies produced in this way could have potential as new ligands for assays of TSH receptor antibodies, TSH and related compounds. Also they may be effective agents in vivo for regulating the action of TSH receptor antibodies, TSH and related compounds.

Other methods of identifying and providing new types of antibody binding sites using monoclonal antibodies are well known. For example by antibody screening of phage-displayed random peptide libraries as described by JC Scott and GP Smith; "Searching for peptide ligands with an epitope library"; Science 1990; 249(4967): 386-390 and MA Myers, JM Davies, JC Tong, J Whisstock, M Scealy, IR MacKay, MJ Rowley; "Conformational epitopes on the diabetes autoantigen GAD<sub>65</sub> identified by peptide phage display and molecular modelling"; Journal of Immunology 2000;

165: 3830-3838. Antibody screening of non-peptide compounds and libraries of non-peptide compounds can also be carried out.

New types of TSH receptor antibody binding sites identified and provided using these procedures may also be useful as new ligands in assays for TSH receptor antibodies, TSH and related compounds. Furthermore they may be effective agents in vivo for regulating the action of TSH receptor antibodies, TSH and related compounds.

In view of the potential value of human monoclonal TSH receptor antibodies there have been considerable efforts over many years to produce such antibodies (see for example B Rees Smith, SM McLachlan, J Furmaniak; "Autoantibodies to the thyrotropin receptor"; Endocrine Reviews 1988; 9: 106-121. However, to date these efforts have been unsuccessful (see for example SM McLachlan, B Rapoport; "Monoclonal, human autoantibodies to the TSH receptor – The Holy Grail and why are we looking for it"; Journal of Clinical Endocrinology and Metabolism 1996; 81: 3152-3154 and JHW van der Heijden, TWA de Bruin, KAFM Gludemans, J de Kruif, JP Banga, T Logtenberg; "Limitations of the semisynthetic library approach for obtaining human monoclonal autoantibodies to the thyrotropin receptor of Graves' disease"; Clinical and Experimental Immunology 1999; 118: 205-212).

It is an object of the present invention to provide a binding partner for the TSH receptor capable of interacting with the TSH receptor in a manner comparable to the interaction of TSH receptor autoantibodies with the TSH receptor, in particular it is an object of the present invention to provide human monoclonal antibodies to the TSH receptor exhibiting a comparable interaction therewith as seen with TSH receptor antibodies present in the sera of patients with hyperthyroid Graves' disease. The considerable difficulties of producing human monoclonal TSH receptor antibodies have been overcome in the invention described herein. In particular the successful production of a human monoclonal TSH receptor antibody with the characteristics of the autoantibodies found in the sera of patients with hyperthyroid Graves' disease is described. The human TSH receptor monoclonal antibody we have produced (described herein as hMAb TSHR 1) binds to the TSH receptor with high affinity and in such a way that small amounts of the antibody inhibit labelled TSH binding to the TSH receptor and small amounts act as powerful thyroid stimulators. Fab fragments

of the antibody are similarly effective thyroid stimulators and inhibitors of labelled TSH binding as intact IgG. Monoclonal Fab and / or intact IgG can be labelled with <sup>125</sup>I or biotin and shown to bind to the TSH receptor. Such binding is inhibited by TSH receptor autoantibodies in patient sera.

There is provided by the present invention, therefore, a binding partner for the TSH receptor, which binding partner comprises, or is derived from, a human monoclonal or recombinant antibody, or one or more fragments thereof, reactive with the TSH receptor.

In particular, there is provided by the present invention a binding partner for the TSH receptor, which binding partner comprises, or is derived from, a human monoclonal antibody, or one or more fragments thereof, reactive with the TSH receptor.

In particular, there is provided by the present invention a human monoclonal antibody, or one or more fragments thereof, reactive with the TSH receptor.

A binding partner according to the present invention, and in particular, a human monoclonal or recombinant antibody reactive with the TSH receptor according to the present invention can be further characterised by its ability to inhibit TSH binding to the TSH receptor, and / or its ability to stimulate the TSH receptor, both of which have been seen to be comparable to the respective inhibitory and stimulatory properties of TSH receptor autoantibodies present in sera obtained from patients with Graves' disease.

More particularly, a binding partner according to the present invention, and in particular a human monoclonal or recombinant antibody according to the present invention, can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments of such a monoclonal or recombinant antibody.

More particularly, a binding partner according to the present invention, and in particular a human monoclonal or recombinant antibody according to the present invention, can be further characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments of such a monoclonal or recombinant antibody.

In a preferred embodiment of the present invention, a binding partner according to the present invention, and in particular a human monoclonal or recombinant antibody according to the present invention, can be characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments of such a monoclonal or recombinant antibody.

In the case where a binding partner according to the present invention comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, in particular for example one or more Fab fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, it may be preferred that such a binding partner can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg.

It may also be preferred in the case where a binding partner according to the present invention comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, in particular for example one or more Fab fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, that such a binding partner can be characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 100 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 200 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 400 units of International Standard NIBSC 90/672 per mg, per mg.

It may be still further preferred in the case where a binding partner according to the present invention comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, in particular for example one or more Fab fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, that such a binding partner can be characterised by:

(i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg; and

(ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 100 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 200 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 400 units of International Standard NIBSC 90/672 per mg.

In a preferred case the present invention provides a binding partner for the TSH receptor (typically a human monoclonal antibody), which binding partner is capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor and which comprises an antibody V<sub>H</sub> domain selected from the group consisting of a V<sub>H</sub> domain as shown in SEQ ID NO. 1 and a V<sub>H</sub> domain comprising one or more V<sub>H</sub> CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4.

In a first embodiment of the present invention, there is, therefore, provided a binding partner for the TSH receptor (typically a human monoclonal antibody), which binding partner is capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor and which comprises an antibody V<sub>H</sub> domain as shown in SEQ ID NO. 1.

In a second embodiment of the present invention there is, therefore, provided a binding partner for the TSH receptor (typically a human monoclonal antibody), which binding partner is capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor and which comprises an antibody V<sub>H</sub> domain comprising one or more V<sub>H</sub> CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4.

It will be appreciated that a binding partner according to the present invention can comprise an antibody  $V_H$  domain substantially as hereinbefore described in the absence of an antibody  $V_L$  domain. It is known that single immunoglobulin domains, especially  $V_H$  domains, are capable of binding target antigens in a specific manner. Alternatively, a binding partner according to the present invention can comprise an antibody  $V_H$  domain paired with an antibody  $V_L$  domain to provide an antibody

binding site comprising both  $V_H$  and  $V_L$  domains for a TSH receptor employing techniques well known in the art (Biochim. Biophys. Acta, 192 (1969) 277-285; Proc. Natl. Acad. Sci. USA, Vol. 89, pp 10026-10030, November 1992).

In a preferred case the present invention provides, however, a binding partner for the TSH receptor, which binding partner is capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor and which comprises:

an antibody V<sub>H</sub> domain selected from the group consisting of:

a  $V_H$  domain as shown in SEQ ID NO. 1 and a  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4; and / or

an antibody V<sub>L</sub> domain selected from the group consisting of:

a  $V_L$  domain as shown in SEQ ID NO. 6 and a  $V_L$  domain comprising one or more  $V_L$  CDRs with an amino acid sequence selected from SEQ ID NO. 7, SEQ ID NO. 8 and SEQ ID NO. 9.

It may be preferred according to the present invention that a binding partner substantially as hereinbefore described comprises an antibody  $V_H$  domain substantially as hereinbefore described paired with an antibody  $V_L$  domain substantially as hereinbefore described to provide an antibody binding site comprising both  $V_H$  and  $V_L$  domains for the TSH receptor, although as discussed further an antibody  $V_H$  domain, or an antibody  $V_L$  domain, may be independently used to bind a TSH receptor. It will be appreciated, therefore, that a binding partner substantially as hereinbefore described can comprise an antibody  $V_H$  domain. It will also be appreciated, therefore, that a binding partner substantially as hereinbefore described in the absence of an antibody  $V_L$  domain. It will also be appreciated, therefore, that a binding partner substantially as hereinbefore described can comprise an antibody  $V_L$  domain. Alternatively, a binding partner substantially as hereinbefore described can comprise an antibody  $V_H$  domain. Alternatively, a binding partner substantially as hereinbefore described can comprise an antibody  $V_H$  domain paired with an antibody

 $V_L$  domain substantially as hereinbefore described to provide an antibody binding site comprising both  $V_H$  and  $V_L$  domains for the TSH receptor.

Preferred embodiments according to the present invention can thus include a binding partner substantially as hereinbefore described comprising an antibody  $V_H$  domain as shown in SEQ ID NO. 1 paired with an antibody  $V_L$  domain as shown in SEQ ID NO. 6 to provide an antibody binding site, comprising both these  $V_H$  and  $V_L$  domains for the TSH receptor.

It is further envisaged according to the present invention that  $V_H$  domains substantially as hereinbefore described may be paired with  $V_L$  domains other than those specifically described herein. It is also further envisaged according to the present invention that  $V_L$  domains substantially as hereinbefore described may be paired with  $V_H$  domains other than those specifically described herein.

According to a further embodiment of the present invention there is provided a binding partner substantially as hereinbefore described for the TSH receptor, which binding partner is capable of binding to the TSH receptor so as to stimulate the TSH receptor and which can comprise:

an antibody V<sub>H</sub> domain comprising:

a  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4; and / or

an antibody V<sub>L</sub> domain comprising:

a  $V_L$  domain comprising one or more  $V_L$  CDRs with an amino acid sequence selected from SEQ ID NO. 7, SEQ ID NO. 8 and SEQ ID NO. 9.

One or more CDRs as referred to above may be taken from the hereinbefore described  $V_{\rm H}$  and  $V_{\rm L}$  domains and incorporated into a suitable framework. For example, the

amino acid sequence of one or more CDRs substantially as hereinbefore described may be incorporated into framework regions of antibodies differing from hMAb TSHR 1 specifically disclosed herein, such antibodies thereby incorporating the one or more CDRs and being capable of binding to the TSH receptor, preferably to stimulate the TSH receptor substantially as hereinbefore described. Alternatively, the present invention may provide a polypeptide capable of binding to the TSH receptor so as to stimulate the TSH receptor substantially as hereinbefore described and comprising the primary structural conformation of amino acids as represented by one or more CDRs as specifically described herein, optionally together with further amino acids, which further amino acids may enhance the binding affinity of one or more CDRs as described herein for the TSH receptor or may have substantially no role in affecting the binding properties of the polypeptide for the TSH receptor.

The present invention, also encompasses variants, analogs, derivatives and fragments of the specific human monoclonal antibody described herein, V<sub>H</sub> domains, CDRs and polypeptides disclosed herein, which variants, analogs, derivatives and fragments retain the ability to interact with the TSH receptor (such as for example to stimulate the TSH receptor) substantially as hereinbefore described.

The terms "variants", "analogs", "derivatives" and "fragments" as used herein can be characterised as antibodies, antibody fragments or polypeptides which retain essentially the same biological function or activity as a human monoclonal antibody having a  $V_H$  domain as shown in SEQ ID NO.1 and a  $V_L$  domain as shown in SEQ ID NO.6 and in particular in respect of the binding properties thereof for the TSH receptor. Suitably, variants, analogs, derivatives and fragments, and variants, analogs and derivatives of the fragments as described herein, have a primary structural conformation of amino acids in which several or a few (such as 5 to 10, 1 to 5 or 1 to 3) amino acid residues of a human monoclonal antibody having a  $V_H$  domain as shown in SEQ ID NO.1 and a  $V_L$  domain as shown in SEQ ID NO.6 are substituted, deleted or added, in any combination. Especially preferred among these are silent substitutions, additions and deletions which do not alter or substantially alter the biological activity or function of a human monoclonal antibody having a  $V_H$  domain as shown in SEQ ID NO.1 and a  $V_L$  domain as shown in SEQ ID NO.6. Conservative substitutions can be preferred as hereinafter described in greater detail.

More particularly, variants, analogs or derivatives of a human monoclonal antibody having a  $V_H$  domain as shown in SEQ ID NO.1 and a  $V_L$  domain as shown in SEQ ID NO.6 according to the present invention may be ones in which one or more of the amino acid residues are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue), or ones in which one or more of the amino acid resides includes a substituent group or the like. Such variants, derivatives and analogs are deemed to be within the scope of those skilled in the art from the teachings herein.

Most typically, variants, analogs or derivatives are those that vary from a reference human monoclonal antibody having a  $V_H$  domain as shown in SEQ ID NO.1 and a  $V_L$  domain as shown in SEQ ID NO.6 by conservative amino acid substitutions. Such substitutions are those that substitute a given amino acid by another amino acid of like characteristics. Typically seen as conservative substitutions are the replacements, one for another, among the aliphatic amino acids A, V, L and I; among the hydroxyl residues S and T; among the acidic residues D and E; among the amide residues N and Q; among the basic residues K and R; and among the aromatic residues F and Y.

It will be appreciated that the term fragment as used herein in particular relates to fragments of antibodies specifically as herein described and form an important aspect of the present invention. In this way, a human monoclonal or recombinant antibody as provided by the present invention may be provided as any of the following fragments: (i) the Fab fragment consisting of V<sub>L</sub>, V<sub>H</sub>, C<sub>L</sub> and C<sub>H</sub>1 domains; (ii) the Fd fragment consisting of the V<sub>H</sub> and C<sub>H</sub>1 domains; (iii) the Fv fragment consisting of the V<sub>L</sub> and V<sub>H</sub> domains; (iv) the dAb fragment which consists of a V<sub>H</sub> domain; (v) isolated CDR regions; (vi) F(ab')2 fragments, a bivalent fragment comprising two linked Fab fragments; and (vii) single chain Fv molecules (scFv), wherein a V<sub>H</sub> domain and a V<sub>L</sub> domain are linked by a peptide linker which allows the two domains to associate to form an antigen binding site.

Alternatively, a human monoclonal or recombinant antibody according to the present invention may comprise a whole IgG antibody, whereby the antibody includes variable and constant regions.

The present invention also provides a further binding partner capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor substantially as hereinbefore described, and which further binding partner can compete for binding to the TSH receptor with a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described, which further binding partner does not comprise TSH or a human antibody to the TSH receptor. The present invention also provides a further binding partner capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor substantially as hereinbefore described, and which further binding partner can compete for binding to the TSH receptor with a binding partner for the TSH receptor comprising a V<sub>H</sub> domain and / or a V<sub>L</sub> domain substantially as hereinbefore described, which further binding partner comprises, or is derived from, a human monoclonal or recombinant antibody, or one or more fragments thereof, reactive with the TSH receptor. In particular this further binding partner may comprise a further antibody having a binding site for an epitope region of the TSH receptor, which further antibody is capable of binding to the TSH receptor preferably so as to stimulate the TSH receptor substantially as hereinbefore described and can compete for binding to the TSH receptor with a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described.

Suitably such a further binding partner can be derived from a specific binding partner as described herein, hMAb TSHR 1, by suitable mutagenesis techniques, such as spot mutations or the like, so as to obtain a further binding partner for the TSH receptor that can compete with a binding partner substantially as herein described (such as hMAb TSHR 1) for interaction with the TSH receptor.

Preferably such a further binding partner for the TSH receptor can comprise a monoclonal or recombinant antibody and can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or

more fragments of the antibody. It may also be preferred that such a further binding partner according to the present invention, can be characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments of the antibody.

It may also be even more preferred that such a further binding partner of the present invention, can be characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

There is also provided by the present invention a polynucleotide comprising:

(i) a nucleotide sequence as shown in SEQ ID NO. 10, SEQ ID NO. 11, SEQ ID NO. 12, SEQ ID NO. 13, SEQ ID NO. 15, SEQ ID NO. 16, SEQ ID NO. 17 or SEQ ID NO. 18, encoding an amino acid sequence of an antibody V<sub>H</sub> domain, V<sub>L</sub> domain, or CDR, as shown in SEQ ID NO. 1, SEQ ID NO. 2,

SEQ ID NO. 3, SEQ ID NO. 4, SEQ ID NO. 6, SEQ ID NO. 7, SEQ ID NO. 8 or SEQ ID NO. 9;

- (ii) a nucleotide sequence encoding a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described, or encoding an amino acid sequence of an antibody  $V_H$  domain,  $V_L$  domain, or CDR, of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described;
- (iii) a nucleotide sequence differing from any sequence of (i) in codon sequence due to the degeneracy of the genetic code;
- (iv) a nucleotide sequence comprising an allelic variation of any sequence of (i);
- (v) a nucleotide sequence comprising a fragment of any of the sequences of (i), (ii), (iii), or (iv) and in particular a nucleotide sequence comprising a fragment of any of the sequences of (i), (ii), (iii), (iv) or (v) and encoding a Fab fragment, a Fd fragment, a Fv fragment, a dAb fragment, an isolated CDR region, F(ab')2 fragments or a scFv fragment, of a human monoclonal antibody substantially as hereinbefore described;
- (vi) a nucleotide sequence differing from the any sequence of (i) due to mutation, deletion or substitution of a nucleotide base and encoding a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described, or encoding an amino acid sequence of an antibody V<sub>H</sub> domain, V<sub>L</sub> domain, or CDR, of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described.

Variant polynucleotides according to the present invention are suitably at least 70% identical over their entire length to any polynucleotide sequence of (i), most highly preferred are polynucleotides that comprise a region that is at least 80% identical over its entire length to any polynucleotide sequence of (i), polynucleotides at least 90%

identical over their entire length to any polynucleotide sequence of (i) are particularly preferred, and among these particularly preferred polynucleotides, those with at least 95% identity are especially preferred.

The present invention further provides a biologically functional vector system which carries a polynucleotide substantially as hereinbefore described and which is capable of introducing the polynucleotide into the genome of a host organism.

The present invention also relates to host cells which are transformed with polynucleotides of the invention and the production of binding partners for the TSH receptor (typically human monoclonal antibodies) of the invention by recombinant techniques. Host cells can be genetically engineered to incorporate polynucleotides and express binding partners for the TSH receptor (typically human monoclonal antibodies) of the present invention.

The amino acid sequences of hMAb TSHR 1, a human monoclonal antibody according to the present invention, and nucleotide sequences coding therefor, are shown in the Sequence listings as herein after described and can be assigned as follows:

### Amino Acid Sequences

SEQ 1D NO. 1	$V_{H}$
SEQ ID NO. 2	V <sub>H</sub> CDRI
SEQ ID NO. 3	V <sub>H</sub> CDRII
SEQ ID NO. 4	V <sub>H</sub> CDRIII
SEQ ID NO. 5	Heavy chain variable and adjacent constant region
SEQ 1D NO. 6	$V_L$
SEQ 1D NO. 7	$V_L$ CDRI
SEQ 1D NO. 8	$V_L CDRII$
SEQ 1D NO. 9	V <sub>L</sub> CDRIII

### Nucleotide Sequences

SEQ ID NO. 10	$V_{H}$
SEQ ID NO. 11	V <sub>H</sub> CDRI
SEQ ID NO. 12	V <sub>H</sub> CDRII
SEQ ID NO. 13	V <sub>H</sub> CDRIII.
SEQ ID NO. 14	Heavy chain variable and adjacent constant region
SEQ ID NO. 15	$V_L$
SEQ ID NO. 16	V <sub>L</sub> CDRI
SEQ ID NO. 17	V <sub>L</sub> CDRII
SEQ ID NO. 18	V <sub>L</sub> CDRIII

The above sequences for TSHR1 can also be seen by reference to Figures 4, 5, 6 and 7.

The present invention also provides a process of providing a human monoclonal antibody to the TSH receptor substantially as hereinbefore described, which process comprises:

- (i) providing a source of lymphocytes from a subject, which subject has TSH receptor antibody activity of greater than about 0.04 units of NIBSC 90/672 per mL of serum with respect to inhibition of TSH binding to the TSH receptor;
- (ii) isolating lymphocytes from said lymphocyte source of (i);
- (iii) immortalising the isolated lymphocytes; and
- (iv) cloning the immortalised lymphocytes so as to produce an immortalised colony secreting a human monoclonal antibody to the TSH receptor substantially as hereinbefore described.

Alternatively, a process of providing a human monoclonal antibody to the TSH receptor substantially as hereinbefore described can be defined as a process which comprises:

- (i) providing a source of lymphocytes from a subject, which subject has TSH receptor antibody activity of greater than about 0.1 units of NIBSC 90/672 per mL of serum with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor;
- (ii) isolating lymphocytes from said lymphocyte source of (i);
- (iii) immortalising the isolated lymphocytes; and
- (iv) cloning the immortalised lymphocytes so as to produce an immortalised colony secreting a human monoclonal antibody to the TSH receptor substantially as hereinbefore described.

Preferably a process according to the present invention comprises isolating lymphocytes from peripheral blood, thyroid tissue, spleen tissue, lymph nodes or bone marrow, most typically from peripheral blood. Typically, the source of lymphocytes for use in a method according to the present invention can be further characterised as being obtained from a subject having serum TSH receptor antibody levels of greater than about 0.1 units of NIBSC 90/672 per mL with respect to inhibition of TSH binding to the TSH receptor, or more typically greater than about 0.2 units of NIBSC 90/672 per mL with respect to inhibition of TSH binding to the TSH receptor, or more typically greater than about 0.3 units of NIBSC 90/672 per mL with respect to inhibition of TSH binding to the TSH receptor and preferably being in the range of about 0.3 to 0.5 units of NIBSC 90/672 per mL or greater with respect to inhibition of TSH binding to the TSH receptor. Alternatively, or additionally, the source of lymphocytes for use in a method according to the present invention can typically be further characterised as being obtained from a subject having serum TSH receptor antibody levels of greater than about 0.2 units of NIBSC 90/672 per mL with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor, or more typically greater than about 0.5 units of NIBSC 90/672 per mL with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor and preferably being in the range of about 0.5 to 1.0 units of NIBSC 90/672 per mL or greater with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor. It will be appreciated from the above that the immune response to the TSH receptor of a subject from which lymphocytes are isolated should preferably be in a highly active phase.

Preferably a process according to the present invention comprises infecting the isolated lymphocytes with Epstein Barr virus, and suitably the thus immortalised lymphocytes are fused with a mouse / human cell line. Suitably a process according to the present invention further comprises screening the resulting clones for TSH receptor antibodies, for example by inhibition of <sup>125</sup>I TSH binding to the TSH receptor in an assay system which has a sensitivity of at least about lunit/L of NIBSC 90/672.

The present invention further provides a human monoclonal antibody to the TSH receptor obtained by a process substantially as described above. Preferably such an obtained human monoclonal antibody to the TSH receptor according to the present invention, can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments of such a human monoclonal antibody.

More particularly, it may be preferred that such a human monoclonal antibody according to the present invention, can be further characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments of such a human monoclonal antibody.

In a preferred embodiment of the present invention, such a human monoclonal antibody according to the present invention, can be characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments of such a human monoclonal antibody.

It may also be preferred that one or more fragments of a thus obtained human monoclonal antibody according to the present invention, in particular for example one or more Fab fragments thereof, can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg. It may also be preferred that such one or more fragments can be characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 100 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 200 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 200 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 400 units of International Standard NIBSC 90/672 per mg.

More preferably, such one or more Fab fragments can be characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 100 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 200 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 400 units of International Standard NIBSC 90/672 per mg.

A process substantially as described above may further comprise a further process stage whereby the obtained human monoclonal antibody is subjected to suitable further processing techniques (such as suitable mutagenesis techniques, such as spot mutations or the like), so as to obtain a further binding partner for the TSH receptor that can compete with a binding partner substantially as herein described (such as hMAb TSHR 1) for interaction with the TSH receptor. Such further processing techniques are well known to one of ordinary skill in the art. The present invention further provides a further binding partner to the TSH receptor obtained by such further processing techniques.

Preferably such a further binding partner for the TSH receptor can comprise a monoclonal or recombinant antibody and can be characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof. It may also be preferred that such a further binding partner according to the present invention, can be characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30

units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.

It may also be even more preferred that such a further binding partner of the present invention, can be characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, or more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 60 units of International Standard NIBSC 90/672 per mg, more preferably of at least about 120 units of International Standard NIBSC 90/672 per mg, or even more preferably of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

A binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention may have diagnostic and therapeutic applications.

Accordingly, a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can be employed in screening methods for detecting autoantibodies to the TSH receptor in patient sera and also in diagnostic methods. In this way, a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can be employed in place of, or in addition to, competitors hitherto described for use in screening methods for

detecting autoantibodies to the TSH receptor and also in diagnostic methods. Similarly, a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can be employed in place of, or in addition to, competitors hitherto described for use in kits for use in detecting autoantibodies to the TSH receptor.

The present invention also provides, therefore, a method of screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to a TSH receptor, said method comprising:

- (a) providing said sample of body fluid from said subject;
- (b) providing one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;
- (c) contacting said sample with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) autoantibodies to the TSH receptor present in said sample, or (ii) said binding partner for the TSH receptor (typically a human monoclonal antibody); and
- (d) monitoring the interaction of said second molecule of said binding pair with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.

It will be appreciated that binding molecules of the one or more binding pairs can be antigen-antibody (for example, [TSH receptor or epitope]—[monoclonal TSH receptor antibody]), anti-idiotypic antibody-monoclonal TSH receptor antibody or novel TSH receptor antibody binding member-monoclonal TSH receptor antibody. Preferably,

the binding molecules of the binding pairs are antigen-antibody, namely, [TSH receptor or one or more epitopes thereof]—[monoclonal TSH receptor antibody], where the epitopes may be "free standing" or present in a larger scaffold polypeptide or the like.

Preferably, the present invention provides, therefore, a method of screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to a TSH receptor, said method comprising:

- (a) providing said sample of body fluid from said subject;
- (b) contacting said sample with (i) a full length TSH receptor, or one or more epitopes thereof or a polypeptide comprising one or more epitopes of a TSH receptor, and (ii) a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention, under conditions that allow interaction of the TSH receptor with autoantibodies produced in response to the TSH receptor, so as to permit said TSH receptor, or said one or more epitopes thereof or said polypeptide, to interact with either autoantibodies to the TSH receptor present in said sample, or said binding partner for the TSH receptor (typically a human monoclonal antibody); and
- (c) monitoring the interaction of said TSH receptor, or said one or more epitopes thereof or said polypeptide, with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.

In certain embodiments, a method according to the present invention may also employ one or more competitors that can compete in the interaction of the binding partner of the present invention and the second molecule of the binding pair, such as the TSH receptor or epitopes thereof substantially as hereinbefore described. Such competitors may comprise TSH, or one or more monoclonals reactive with the TSH receptor, such as mouse monoclonals reactive with the TSH receptor.

Preferably, a method according to the present invention as referred to above, further comprises providing labelling means for a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and where appropriate one or more competitors as described above, suitable labelling means including enzymic labels, isotopic labels, chemiluminescent labels, fluorescent labels, dyes and the like.

The present invention also provides, a kit for screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to a TSH receptor, said kit comprising:

- (a) one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;
- (b) means for contacting said sample of body fluid from said subject with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) autoantibodies to the TSH receptor present in said sample, or (ii) said binding partner for the TSH receptor (typically a human monoclonal antibody); and
- (c) means for monitoring the interaction of said second molecule of said binding pair with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.

It will be appreciated that binding molecules of the one or more binding pairs can be antigen-antibody (for example, [TSH receptor or epitope]—[monoclonal TSH receptor antibody]), anti-idiotypic antibody-monoclonal TSH receptor antibody or novel TSH receptor antibody binding member-monoclonal TSH receptor antibody. Preferably,

the binding molecules of the binding pairs are antigen-antibody, namely, [TSH receptor or one or more epitopes thereof]—[monoclonal TSH receptor antibody], where the epitopes may be "free standing" or present in a larger scaffold polypeptide or the like.

The present invention preferably provides a kit for screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to the TSH receptor, said kit comprising:

- (a) a full length TSH receptor, or one or more epitopes thereof or a polypeptide comprising one or more epitopes of the TSH receptor;
- (b) a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention;
- (c) means for contacting said sample of body fluid from said subject, said TSH receptor, or said one or more epitopes thereof or said polypeptide, and said binding partner for the TSH receptor (typically a human monoclonal antibody), under conditions that allow interaction of the TSH receptor with autoantibodies produced in response to the TSH receptor, so as to permit said TSH receptor, or said one or more epitopes thereof or said polypeptide, to interact with either autoantibodies to a TSH receptor present in said sample, or said binding partner for the TSH receptor (typically a human monoclonal antibody); and
- (d) means for monitoring the interaction of said TSH receptor, or said one or more epitopes thereof or said polypeptide, with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.

In certain embodiments, a kit according to the present invention may further comprise one or more competitors that can compete in the interaction of the binding partner of the present invention and the second molecule of the binding pair, such as the TSH receptor or epitopes thereof substantially as hereinbefore described. Such competitors may comprise TSH, or one or more monoclonals reactive with the TSH receptor, such as mouse monoclonals reactive with the TSH receptor.

Suitably, a kit as referred to above further comprises labelling means for a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and where appropriate one or more competitors as described above, suitable labelling means being substantially as hereinbefore described.

In the presence of autoantibodies to the TSH receptor, binding of the TSH receptor to a binding partner for the TSH receptor (typically a human monoclonal antibody) in a method or kit as described above will be decreased.

A binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can also be employed in assay methods and kits substantially as described above for TSH and related ligands.

The present invention also provides, therefore, a method of assaying TSH and related ligands, said method comprising:

- (a) providing a sample suspected of containing or containing TSH or related ligands;
- (b) providing one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;
- (c) contacting said sample with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) TSH or related ligands present in said sample, or (ii) said binding partner for the TSH receptor (typically a human monoclonal antibody); and

(d) monitoring the interaction of said second molecule of said binding pair with TSH or related ligands present in said sample, thereby providing an indication of the presence of TSH or related ligands in said sample.

The present invention also provides a kit for assaying TSH or related ligands, said kit comprising:

- (a) one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;
- (b) means for contacting a sample suspected of containing or containing TSH or related ligands with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) TSH or related ligands present in said sample, or (ii) said binding partner for the TSH receptor (typically a human monoclonal antibody); and
- (c) means for monitoring the interaction of said second molecule of said binding pair with TSH or related ligands present in said sample, thereby providing an indication of the presence of TSH or related ligands in said sample.

A further application of a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention is its use to identify and provide new types of TSH receptor antibody binding sites. There is further provided by the present invention, therefore, a process of identifying one or more epitope regions of the TSH receptor, which process comprises contacting a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described with a full length TSH receptor, or one or more fragments thereof, so as to allow interaction of said binding partner for the TSH receptor with said full length TSH receptor, or said one or more fragments thereof, and identifying the amino acids of said full length TSH receptor, or said one or more fragments

thereof, with which said binding partner interacts. Suitably, interaction of the binding partner with selected fragments of the TSH receptor and the full length TSH receptor, is analysed, so as to identify the amino acids of the TSH receptor with which the binding partner interacts.

Furthermore, the present invention allows for generation of antibodies to the regions of a monoclonal TSH receptor antibody according to the present invention which bind the TSH receptor. Such anti-idiotypic antibodies produced in this way could have potential as new ligands for assays of TSH receptor autoantibodies, TSH and related compounds. Also they may be effective agents in vivo for regulating the action of TSH receptor autoantibodies, TSH and related compounds. The present invention further provides, therefore, one or more anti-idiotypic antibodies generated to binding regions of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described.

Other methods of identifying and providing new types of antibody binding sites using monoclonal antibodies are well known. For example by antibody screening of phage-displayed random peptide libraries as described by JC Scott and GP Smith; "Searching for peptide ligands with an epitope library"; Science 1990; 249(4967): 386-390 and MA Myers, JM Davies, JC Tong, J Whisstock, M Scealy, IR MacKay, MJ Rowley; "Conformational epitopes on the diabetes autoantigen GAD<sub>65</sub> identified by peptide phage display and molecular modelling"; Journal of Immunology 2000; 165: 3830-3838. Antibody screening of non-peptide compounds and libraries of non-peptide compounds can also be carried out.

New types of TSH receptor antibody binding sites identified and provided using these procedures may also be useful as new ligands in assays for TSH receptor autoantibodies, TSH and related compounds. Furthermore they may be effective agents in vivo for regulating the action of TSH receptor autoantibodies, TSH and related compounds.

A binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention substantially as hereinbefore described can also be usefully employed in therapy. There is, therefore, further provided by the present

invention methods of treatment comprising administration of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described, pharmaceutical compositions comprising a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described (together with one or more pharmaceutically acceptable carriers, diluents or excipients therefor), and use of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described in the manufacture of a medicament or composition.

A binding partner for the TSH receptor, in particular a human monoclonal antibody to the TSH receptor derived from patients' lymphocytes according to the present invention, is a valuable reagent for understanding the pathogenesis of Graves' disease and for developing new methods of measuring TSH receptor autoantibodies, for example as replacements for TSH in competitive binding assays substantially as hereinbefore described. Also, a stimulating binding partner according to the present invention has in vivo applications when tissue containing the TSH receptor (eg thyroid tissue or thyroid cancer tissue) requires stimulation. The present invention provides, therefore, a medicament or composition for use in stimulating thyroid tissue, and / or tissue containing the TSH receptor. In particular, a stimulating binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can be employed in oncology, and in particular for use in the diagnosis, management and treatment of thyroid cancer. Alternatively, a binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can be a powerful TSH antagonist (blocking antibody) and such a monoclonal TSH receptor antibody according to the present invention being a TSH receptor antagonist is valuable for in vivo applications when the activity of tissue containing the TSH receptor (eg thyroid tissue or thyroid cancer tissue) requires inactivation or to be made unresponsive to TSH, TSH receptor autoantibodies or other stimulators.

One of the major advantages of a monoclonal antibody as provided by the present invention over TSH in such in vitro and / or in vivo applications is the relative ease with which such antibodies can be manipulated. For example, manipulation of the TSH receptor binding region of a monoclonal antibody according to the present

invention so as to change the characteristics thereof, such as affinity and biological characteristics, including the degree of TSH agonist or antagonist activities. Also monoclonal antibodies according to the present invention have a much longer half life than TSH in vivo and this may have considerable advantages in in vivo applications. Furthermore, the half life of the antibodies can be manipulated, for example antibody Fab fragments have a much shorter half life than intact IgG.

Pharmaceutical compositions according to the present invention include those suitable for oral, parenteral and topical administration, although the most suitable route will generally depend upon the condition of a patient and the specific disease being treated. The precise amount of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described to be administered to a patient will be the responsibility of an attendant physician, although the dose employed will depend upon a number of factors, including the age and sex of the patient, the specific disease being treated and the route of administration substantially as described above.

There is further provided by the present invention a method of stimulating thyroid tissue, and / or tissue containing a TSH receptor, which method comprises administering to a patient in need of such stimulation a diagnostically or therapeutically effective amount of a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described.

The present invention also provides in combination, a binding partner for the TSH receptor (typically a human monoclonal antibody) substantially as hereinbefore described, together with one or more further agents capable of stimulating thyroid tissue, and / or tissue containing a TSH receptor, for simultaneous, separate or sequential use in stimulating thyroid tissue, and / or tissue containing a TSH receptor. Preferably the one or more further agents comprise recombinant human TSH and / or one or more variants, analogs, derivatives or fragments thereof, or variants, analogs or derivatives of such fragments. Alternatively, the one or more further agents can act independently of binding to the TSH receptor.

A binding partner for the TSH receptor (typically a human monoclonal antibody) according to the present invention can also be employed as a replacement source for patient serum required to contain TSH receptor antibody or antibodies for use in commercial kits. Furthermore, a binding partner for the TSH receptor (typically a human monoclonal antibody) can be provided according to the present invention in a preparation required to comprise a defined concentration of TSH receptor antibody or antibodies, and in this way there can be provided a preparation with a defined activity, such as stimulatory activity, with respect to the TSH receptor. Optionally, such a preparation may further comprise one or more further human monoclonal antibodies, such as monoclonal antibodies to GAD, TPO or the like.

The following illustrative explanations are provided to facilitate understanding of certain terms used herein. The explanations are provided as a convenience and are not limitative of the invention

BINDING PARTNER FOR A TSH RECEPTOR, describes a molecule having a binding specificity for the TSH receptor. A binding partner as described herein may be naturally derived or wholly or partially synthetically produced. Such a binding partner has a domain or region which specifically binds to and is therefore complementary to one or more epitope regions of the TSH receptor. In particular, a binding partner as described herein can be a monoclonal antibody to the TSH receptor, and more particularly can be a human monoclonal antibody to the TSH receptor.

C DOMAIN denotes a region of relatively constant amino acid sequence in antibody molecules.

CDR denotes complementarity determining regions which are present on both heavy and light chains of antibody molecules and represent regions of most sequence variability. CDRs represent approximately 15 to 20% of variable domains and represent antigen binding sites of an antibody.

FR denotes framework regions and represent the remainder of the variable light domains and variable heavy domains not present in CDRs.

HC denotes part of a heavy chain of an antibody molecule comprising the heavy chain variable domain and the first domain of an IgG constant region.

HOST CELL is a cell which has been transformed or transfected, or is capable of transformation or transfection by an exogenous polynucleotide sequence.

IDENTITY, as known in the art, is the relationship between two or more polypeptide sequences, or two or more polynucleotide sequences, as determined by comparing the sequences.

LC denotes a light chain of an antibody molecule.

NIBSC 90/672 is an International Standard for thyroid stimulating antibody. The International Standard for thyroid stimulating activity consists of a batch of ampoules containing freeze dried plasma proteins from a single human patient with high TSH receptor autoantibodies. The preparation has been evaluated in an international collaborative study and shown to possess both thyroid stimulating and thyroid receptor binding activity. At the 46<sup>th</sup> meeting in 1995, the Expert Committee on Biological Standardization of WHO established the preparation coded 90/672 as the International Standard for thyroid stimulating antibody. Each ampoule contains freeze-dried residue of 1.0ml of a solution containing 0.02M phosphate buffer, dialysed human plasma proteins and 0.1 International Units (100 milli-International Units) per ampoule by definition.

STIMULATION OF A TSH RECEPTOR by a human monoclonal antibody as described herein denotes the ability thereof to bind to a TSH receptor and to thereby effect, for example, production of cyclic AMP as a result of such binding to the TSH receptor. Such stimulation is analogous to the responses seen on binding of TSH, or TSH receptor autoantibodies, to the TSH receptor and in this way a human monoclonal antibody as described herein essentially provides the same or similar binding responses as seen with TSH, or TSH receptor autoantibody, binding to a TSH receptor.

V DOMAIN denotes a region of highly variable amino acid sequence in antibody molecules.

V<sub>H</sub>DOMAIN denotes variable regions or domains in heavy chains of antibody molecules.

V<sub>L</sub>DOMAIN denotes variable regions or domains in light chains of antibody molecules.

The present invention will now be illustrated by the following Figures and Examples, which do not limit the scope of the invention in any way.

### **Examples**

### **MATERIALS & METHODS**

# Lymphocyte isolation and cloning of human monoclonal TSH receptor autoantibodies

Blood was obtained from a patient with Graves' disease and Type 1 diabetes mellitus who had high levels of serum autoantibodies to the TSH receptor (TRAb). Ethical Committee approval was obtained for the studies. Peripheral blood lymphocytes were isolated on Ficoll-Paque (Amersham Biosciences; Chalfont St Giles, HP8 4SP, UK) from a 20mL blood sample and then infected with Epstein Barr virus (EBV) (European Collection of Cell Cultures - ECACC; Porton Down, SP4 0JG, UK) and cultured on mouse macrophage feeder layers as described before (N Hayakawa, LDKE Premawardhana, M Powell, M Masuda, C Arnold, J Sanders, M Evans, S Chen, JC Jaume, S Baekkeskov, B Rees Smith, J Furmaniak; "Isolation and characterization of human monoclonal autoantibodies glutamic acid Autoimmunity 2002; 35: 343-355). EBV immortalised B decarboxylase"; lymphocytes were then fused with the mouse/human hybrid cell line K6H6/B5 (WL Carroll, K Thilemans, J Dilley, R Levy; "Mouse x human heterohybridomas as fusion partners with human B cell tumors"; Journal of Immunological Methods 1986; 89: 61-72) and cloned two times by limiting dilution at 5 cells/well and a final time at ½ cell/well to obtain a single colony (BJ Bolton, NK Spurr. "B-lymphocytes" In: RI Freshney, MG Freshney (eds). Culture of immortalized cells. Wiley-Liss, New York 1996; 283-297). The original wells and subsequent clones were screened for TSH receptor autoantibody by inhibition of <sup>125</sup>I-TSH binding to solubilised TSH receptor (see below). The single clones producing TSH receptor autoantibodies were grown up in tissue culture flasks.

# Purification and labelling of monoclonal TSH receptor autoantibody preparations

IgGs were purified from tissue culture supernatants using affinity chromatography on Prosep A (Millipore UK Ltd.; Watford, WD18 8YH, UK) according to the manufacturer's instructions and purity assessed by SDS-polyacrylamide gel electrophoresis (PAGE).

Human heavy chain isotype was determined using a radial diffusion assay (The Binding Site; Birmingham, B29 6AT, UK). Human light chain isotype was determined using Western blotting with anti-human kappa chain and anti human lambda chain specific mouse monoclonal antibodies (Sigma-Aldrich Company Ltd; Gillingham, SP8 4XT, UK).

Mouse TSH receptor MAb IgGs were produced and purified as described before (Y Oda, J Sanders, M Evans, A Kiddie, A Munkley, C James, T Richards, J Wills, J Furmaniak, B Rees Smith; "Epitope analysis of the human thyrotropin (TSH) receptor using monoclonal antibodies"; Thyroid 2000; 10: 1051-1059).

The purified IgG preparations were treated with mercuripapain (Sigma-Aldrich) at an enzyme/protein ratio of between 1:10 and 1:100 (depending on the particular monoclonal antibody) and passed through a Prosep A column to remove any intact IgG or Fc fragment from the Fab preparation (Y Oda, J Sanders, S Roberts, M Maruyama, R Kato, M Perez, VB Petersen, N Wedlock, J Furmaniak, B Rees Smith; "Binding characteristics of antibodies to the TSH receptor"; Journal of Molecular Endocrinology 1998; 20: 233-244). Intact IgG was undetectable by SDS-PAGE in the Fab preparations. IgG and Fab preparations of the monoclonal antibodies were labelled with <sup>125</sup>I as described previously (Y Oda, J Sanders, S Roberts, M Maruyama, R Kato, M Perez, VB Petersen, N Wedlock, J Furmaniak, B Rees Smith; "Binding characteristics of antibodies to the TSH receptor"; Journal of Molecular

Endocrinology; 1998; 20: 233-244). IgG preparations were labelled with biotin hydrazide (Pierce Rockford IL61105 USA) according to the manufacturers instructions.

### **Patients**

Sera from patients with Graves' disease of different disease duration were studied. The patients' sera studied showed inhibition of <sup>125</sup>I-labelled TSH binding to the TSH receptor (see below). In addition, sera from 2 patients with Addison's disease (A1 and A2) and high levels of autoantibodies to 21-OH (113 and 1970 units per mL, RSR kit) and 1 serum from a patient with type 1 diabetes mellitus (D1) with high levels of GAD<sub>65</sub> (3700 units per mL; RSR kit) were studied. Informed consent for the study was obtained from the patients. Sera from healthy blood donors (purchased from Golden West Biologicals, Vista, CA 92083, USA) were also studied. TRAb first international standard preparation (90/672) was obtained from the National Institute for Biological Standards and Control (NIBSC; Potters Bar, EN6 3QH, UK).

# Inhibition of <sup>125</sup>I-TSH binding to the TSH receptor

TSH binding inhibition assays were carried out using TSH receptor coated tubes as described previously (J Sanders, Y Oda, S Roberts, A Kiddie, T Richards, J Bolton, V McGrath, S Walters, D Jaskolski, J Furmaniak, B Rees Smith; "The interaction of TSH receptor autoantibodies with <sup>125</sup>I-labeled TSH receptor"; Journal of Clinical Endocrinology and Metabolism 1999; **84**: 3797-3802) (reagents from RSR Ltd). Briefly, 100μL of sample (tissue culture supernatant, purified IgG or Fab fragment, patient serum or NIBSC 90/672 standards) were incubated in TSH receptor coated tubes at room temperature for 2 hours with gentle shaking. After aspiration, the tubes were washed twice with 1 mL of assay buffer (50 mmol/L NaCl, 10 mmol/L Tris-HCl pH 7.8, 0.1% Triton X-100) before addition of 100μL of <sup>125</sup>I-TSH (80,000 cpm) and incubation at room temperature for 1 hour with shaking. The tubes were then washed twice with 1mL of assay buffer, aspirated and counted in a gamma counter. Inhibition of TSH binding was calculated as:-

100 x 1 - cpm TSH bound in the presence of test material cpm bound in the presence of control material

Control materials used were culture medium, a pool of healthy blood donor sera or as otherwise indicated.

# Analysis of thyroid stimulating activities

The ability of monoclonal autoantibody preparations and patient sera to stimulate the production of cyclic AMP in CHO cells expressing hTSH receptor (approximately 50,000 receptors per cell) (Y Oda, J Sanders, S Roberts, M Maruyama, R Kato, M Perez, VB Petersen, N Wedlock, J Furmaniak, B Rees Smith; "Binding characteristics of antibodies to the TSH receptor"; Journal of Molecular Endocrinology 1998; 20: 233-244) were carried out according to the method of R Latif, P Graves, TF Davies; Journal of Biological "Oligomerization of the human thyrotropin receptor"; Chemistry 2001; 276: 45217-45224. Briefly, CHO cells were seeded into 96 well plates (30,000 cells per well) and incubated for 24 hours in DMEM (Invitrogen Ltd; Paisley PA4 9RF, UK) containing 10% fetal calf serum. Culture was then continued in DMEM without fetal calf serum for a further 24 hours. The DMEM was then removed and test IgG, Fab and serum (diluted in NaCl free Hank's Buffered Salts solution containing 1g/L glucose, 20 mmol/L Hepes, 222 mmol/L sucrose, 15 g/L bovine serum albumin (BSA) and 0.5 mmol/L 3 isobutyl-1-methyl xanthine pH 7.4) added and incubated for 1 hour at 37°C. After removal of the test solutions, cells were lysed and assayed for cyclic AMP using a Biotrak enzyme immunoassay system from Amersham Biosciences; Chalfont St Giles, HP8 4SP, UK

# Variable Region Gene Analysis

Total RNA was prepared from 1 x 10<sup>7</sup> cells of a TSH receptor autoantibody producing clone using the acid phenol guanidine method (P Chomczynski, N Sacchi; "Single-step method of RNA isolation by acid guanidinium thiocyanate-phenol-chloroform extraction"; Analytical Biochemistry 1987; 162: 156-159) and mRNA prepared using oligo dT magnetic beads (Dynal Biotech Ltd; Wirral, CH62 3QL, UK). RT-PCR reactions were performed using reagents from Invitrogen Ltd; Paisley PA4 9RF, UK.

Sense strand oligonucleotide primers were designed using the recommended by the Medical Research Council's V-base database (www.mrccpe.cam.ac.uk). Antisense primers specific for human IgG1 heavy chain and lambda light chain were based on constant region encoding DNA sequences. Both sense and antisense primers included additional 5' restriction endonuclease site sequences to facilitate cloning of PCR products. IgG1 heavy chain and lambda light chain RT-PCR reactions were performed using the complete panel of appropriate primers. All primers were synthesized by Invitrogen Ltd. The RT reaction took place at 50°C for 10 minutes followed immediately by 40 cycles of PCR (15 sec 94°C, 30 sec 55°C, 30 sec 72°C). RT-PCR products were cloned into pUC18 and DNA prepared using the Wizard kit from Promega UK Ltd; Southampton SO16 7NS, UK and sequenced by the Sanger-Coulson method (F Sanger, S Nicklen, AR Coulson; "DNA sequencing with chain terminating inhibitors"; Proceedings of the National Academy of Sciences of the USA 1977; 74: 5463-5467). V region sequences were compared with available sequences of human Ig genes using Ig blast (www.ncbi.nlm.nih.gov/ igblast/igblast.cgi).

## Immunoprecipitation Assay (IPA)

The cDNA encoding full length TSH receptor was placed downstream of the T7 promoter in pYES2 (Invitrogen) and used in an in vitro TnT system (Promega UK Ltd) to produce TSH receptor labelled with <sup>35</sup>S-methionine as previously described (L Prentice, J Sanders, M Perez, R Kato, J Sawicka, Y Oda, D Jaskolski, J Furmaniak, B Rees Smith; "Thyrotropin (TSH) receptor autoantibodies do not appear to bind to the TSH receptor produced in an in vitro transcription/translation system"; Journal of Clinical Endocrinology and Metabolism 1997; 82: 1288-1292). Briefly 50μL <sup>35</sup>S-labelled TSH receptor (25 000 – 30 000 cpm) diluted in HSB (150 mmol/L Tris-HCL pH 8.3, 200 mmol/L NaCl and 10 mg/mL bovine serum albumin containing 1% Tween 20) were added to duplicate 50μL aliquots of diluted test sample and incubated for 2 hours at room temperature. Immune complexes were then precipitated by addition of protein A sepharose (Sigma-Aldrich) and counted in a scintillation counter.

### TSH receptor preparations and western blotting

Full-length human TSH receptor was expressed in CHO-K1 cells, extracted with 1% Triton X-100 and purified by TSH receptor monoclonal antibody affinity chromatography as described previously (Y Oda, J Sanders, M Evans, A Kiddie, A Munkley, C James, T Richards, J Wills, J Furmaniak, B Rees Smith; "Epitope analysis of the human thyrotropin (TSH) receptor using monoclonal antibodies"; Thyroid 2000; 10: 1051-1059).

The purified CHO cell produced TSH receptor was run on 9% SDS-PAGE gels, blotted onto nitrocellulose and reacted with test antibodies as described previously (Y Oda, J Sanders, M Evans, A Kiddie, A Munkley, C James, T Richards, J Wills, J Furmaniak, B Rees Smith; "Epitope analysis of the human thyrotropin (TSH) receptor using monoclonal antibodies"; Thyroid 2000; 10: 1051-1059).

## Epitope analysis using TSH receptor peptides

Twenty six peptides each 25aa long covering the whole of the extracellular domain of the human TSH receptor were kindly provided by Dr J Morris (JC Morris, ER Bergert, DJ McCormick; "Structure-function studies of the human thyrotropin receptor. Inhibition of binding of labeled thyrotropin (TSH) by synthetic human TSH receptor peptides"; Journal of Biological Chemistry 1993; 268: 10900-10905). A human 21-OH peptide (C1, SSSRVPYKDRARLPL) which binds to an M21-OH5 MAb (S Chen, J Sawicka, L Prentice, JF Sanders, H Tanaka, V Petersen, C Betterle, M Volpato, S Roberts, M Powell, B Rees Smith, J Furmaniak; "Analysis of autoantibody epitopes on steroid 21-hydroxylase using a panel of monoclonal antibodies"; Journal of Clinical Endocrinology and Metabolism 1998; 83: 2977-2986) was used as a positive control and a human monoclonal antibody to GAD<sub>65</sub> (N Hayakawa, LDKE Premawardhana, M Powell, M Masuda, C Arnold, J Sanders, M Evans, S Chen, JC Jaume, S Baekkeskov, B Rees Smith, J Furmaniak; "Isolation and characterization of human monoclonal autoantibodies to decarboxylase"; Autoimmunity 2002; 35: 343-355) was used as a negative control. The peptide ELISA was carried out as described previously (Y Oda, J Sanders, M Evans, A Kiddie, A Munkley, C James, T Richards, J Wills, J Furmaniak, B Rees

Smith; "Epitope analysis of the human thyrotropin (TSH) receptor using monoclonal antibodies"; Thyroid 2000; 10: 1051-1059).

Interaction of monoclonal TSHR autoantibody preparations with the TSH receptor coated onto plastic tubes or ELISA plate wells

# (a) 125 I-labelled autoantibody

Test samples including patient sera (100μL) were incubated in TSH receptor coated tubes (RSR Ltd.) at room temperature for 2 hours with gentle shaking. After aspiration, the tubes were washed twice with 1mL of assay buffer before addition of 100μL of labelled autoantibody preparation (30,000 cpm) and incubation at room temperature for 1 hour with shaking. The tubes were then washed twice with 1mL of assay buffer, aspirated and counted in a gamma counter. Inhibition of <sup>125</sup>I-labelled autoantibody binding was calculated using the formula as for inhibition of TSH binding (see above).

# (b) Biotin labelled monoclonal autoantibody and biotin labelled TSH

The procedure described previously (J Bolton, J Sanders, Y Oda, C Chapman, R Konno, J Furmaniak and B Rees Smith; "Measurement of thyroid-stimulating hormone receptor autoantibodies by ELISA; Clinical Chemistry, 1999; 45: 2285-2287) was used. Briefly, test samples including patient sera (75μL) were incubated in TSH receptor coated ELISA plate wells (RSR Ltd) for 2 hours with shaking (200 shakes per minute) on an ELISA plate shaker. Test samples were then removed and the wells washed once with assay buffer. Biotin-labelled monoclonal TSH receptor autoantibody (1 ng in 100μL) or biotin labelled porcine TSH (RSR Ltd; 5 ng in 100μL) were then added and incubation continued for 25 minutes at room temperature without shaking. The wells were washed once, 100μL of streptavidin-peroxidase (RSR Ltd; 10 ng in 100μL) added and incubation continued for 20 minutes at room temperature without shaking. The wells were then washed 3 times, the peroxidase substrate tetramethyl benzidine (RSR Ltd; 100μL) added. After incubation for 30 minutes at room temperature without shaking 50μL of 0.5M H<sub>2</sub>SO<sub>4</sub> was added to stop

the substrate reaction and the absorbance of each well read at 450nm on an ELISA plate reader. Inhibition of biotinylated MAb or TSH binding was expressed as an index calculated as:-

100 x 1 - test sample absorbance at 450nm negative serum control absorbance at 450nm

Scatchard analysis of monoclonal autoantibody binding to TSH receptor coated tubes

Unlabelled IgG or Fab in 50µL of assay buffer and 50µL of <sup>125</sup>I-labelled hMAb IgG or Fab (30,000 cpm in assay buffer) were incubated for 2 hours at room temperature with gentle shaking, washed twice with 1mL of assay buffer and counted in a gamma counter. The concentration of IgG or Fab bound vs bound/free was plotted (G Scatchard; "The attraction of proteins for small molecules and ions"; Annals of the New York Academy of Sciences 1949; 51: 660-672) to derive the association constants.

Binding of TSH receptor to tubes coated with monoclonal TSH receptor autoantibodies

Test samples including patient sera (100μL) and detergent solubilised TSH receptor (20μL) were incubated for 1 hour at room temperature. Duplicate 50μL aliquots of the incubation mixture were then added to plastic tubes (Nunc Maxisorp) which had been coated with monoclonal TSH receptor autoantibody Fab (200μL of 10μg/mL overnight at 4°C followed by washing and post coating). After incubation for 1 hour at room temperature with gentle shaking, the tubes were washed, 100μL (40,000 cpm) of <sup>125</sup>I-labelled TSH receptor C terminal monoclonal antibody 4E31 (J Sanders, Y Oda, A Kiddie, T Richards, J Bolton, V McGrath, S Walters, D Jaskolski, J Furmaniak, B Rees Smith; "The interaction of TSH receptor autoantibodies with <sup>125</sup>I-labelled TSH receptor"; Journal of Clinical Endocrinology and Metabolism 1999; 84: 3797-3802) added and incubation continued for a further 1 hour with gentle shaking. Then tubes were then washed and counted for <sup>125</sup>I.

### RESULTS

Lymphocytes (30 x 10<sup>6</sup>) obtained from 20mL of patient's blood were plated out at 1 x 10<sup>6</sup> per well on a 48 well plate with 200µL of EBV supernatant on feeder layers of mouse macrophages. On day 11 post EBV infection the supernatants were monitored for inhibition of <sup>125</sup>I-TSH binding. One well was found to be positive for inhibition of binding, the levels of inhibition increasing to greater than 90% inhibition by day 16 and stayed at that level until day 24, after which time they decreased. The cultures were expanded and fused with K6H6/B5 cells on day 21, 23, 26 and 27 post EBV infection; in total 7 fusion experiments were carried out. Each fusion was plated across 3 x 96 well plates (ie 21 plates in total) and one well, stably producing antibodies with <sup>125</sup>I-TSH binding inhibiting activity was obtained. This was followed by 3 rounds of re-cloning to yield a single clone producing a human monoclonal antibody which inhibited labelled TSH binding to the TSH receptor. This human monoclonal TSH receptor autoantibody was designated hMAb TSHR1 and was of subclass IgG1 with a lambda light chain.

The ability of different concentrations of hMAb TSHR1 IgG and Fab to inhibit labelled TSH binding to the TSH receptor is shown in Figure 1. As can be seen in Figure 1 as little as 1ng/mL of these preparations inhibited TSH binding with more than 90% inhibition being obtained with 1000ng/mL. TSMAb TSHR1 IgG and Fab also stimulated cyclic AMP production in CHO cells transfected with the TSH receptor as shown in Figure 2. As little as 1ng/mL of hMAb TSHR1 IgG or Fab caused strong stimulation of cyclic AMP. Similar levels of stimulation were observed with 0.1ng/mL porcine TSH and 10ng/mL of human TSH. Comparison of the ability of the serum from the original lymphocyte donor (taken at the same time as the blood sample for lymphocyte isolation) to inhibit labelled TSH binding to the TSH receptor and to stimulate cyclic AMP production in TSH receptor transfected CHO cells is shown in Figure 3. Inhibition of TSH binding could be detected with serum diluted 5000x whereas stimulation of cyclic AMP could be detected with serum diluted 5000x.

<sup>125</sup>I-labelled hMAb TSHR1 IgG bound to TSH receptor coated tubes and Scatchard analysis indicated an association constant of 5 x 10<sup>10</sup> molar<sup>-1</sup>. This binding was inhibited by sera from patients with Graves' disease who had TSH receptor

autoantibodies (detectable by inhibition of labelled TSH binding) (Table 1).  $^{125}$ I-labelled hMAb TSHR1 Fab also bound to TSH receptor coated tubes (association constant by Scatchard analysis =  $4.5 \times 10^{10}$  molar<sup>-1</sup>) and this binding was inhibited by TSH receptor autoantibody positive Graves' sera (Table 2). In addition, detergent solubilised preparations were able to bind to plastic tubes coated with hMAb TSHR1 and this binding could be inhibited by sera containing TSH receptor autoantibodies (Table 3).

As shown in Table 4 hMAb TSHR1-biotin bound to TSH receptor coated ELISA plates and the binding was inhibited by the international reference preparation NIBSC 90/672 and serum from patients with Graves' disease. Inhibition of binding was not observed by sera from healthy blood donors

hMAb TSHR1 IgG did not react with full length TSH receptor preparations on Western blot analysis nor did it react well with <sup>35</sup>S-labelled full length TSH receptor in the immunoprecipitation assay nor in the TSH receptor peptide ELISA. This lack of reactivity indicates that hMAb TSHR1 reacts with conformational rather than linear epitopes on the TSH receptor.

Sequence analysis of the genes coding for hMAb TSHR1 indicated that the heavy chain V region genes were of the VH5 family, the D gene of the D6-13 family and the J gene of the JH5 family and for the light chain the V-gene region is from the VL1-11 germline and the J-gene region is from the JL3b germline. The heavy chain nucleotide and amino acid sequences are shown in Figures 4 and 5 respectively and the light chain nucleotide and amino acid sequences are shown in Figures 6 and 7 respectively.

Comparison of the activities of hMAb TSHR1 IgG preparations and the international standard for TSH receptor autoantibodies in terms of inhibition of labelled TSH binding are shown in Table 5. This enabled a specific activity of hMAb TSHR1 IgG to be estimated as 138 units of NIBSC 90/672 per mg of protein when the assays were carried out in serum and 163 units per mg when the assays were carried out in assay buffer (mean of the 2 values = 150 units/mg). hMAb TSHR1 Fab preparations were 288 and 309 units per mg in serum and assay buffer respectively (mean of the 2

values = 300 units/mg). Table 6 shows a similar analysis of the lymphocyte donor serum and the donor serum IgG. As can be seen the donor serum contains a mean of 0.38 units/mL of NIBSC 90/672 (0.36 and 0.4 in serum and assay buffer respectively) and the donor serum IgG has a mean specific activity of 0.059 units per mg of protein. These results are summarised in Table 7 and comparison with the specific activity of hMAb TSHR1 IgG (150 units/mg) indicates that the monoclonal autoantibody IgG is 2500 times more active than the lymphocyte donor serum IgG in terms of inhibition of TSH binding.

Initial assessment of the activities of the various IgG and serum preparations in terms of stimulation of cyclic AMP in CHO cells transfected with the TSH receptor are also shown in Table 7. The stimulation of cyclic AMP assay is characterized by considerable within assay and between assay variability. This relates to several factors including variation in the number and quality of cells initially seeded into the 96 well plates and variation in growth rate of the seeded cells over the subsequent 48 hours. Consequently the assays of hMAb TSHR1 IgG and Fab, lymphocyte donor serum and serum IgG and NIBSC 90/672 were carried out repeatedly and the results are summarized in Table 8. The specific activity of the hMAb TSHR1 IgG was 318 units per mg in the stimulation assay compared with 0.1 units per mg for the lymphocyte donor serum IgG ie the monoclonal autoantibody IgG was about 3000 times as active as the donor serum IgG in terms of stimulation of cyclic AMP production. This value is in reasonable agreement with the value of 2500 times observed for inhibition of TSH binding measurements (see above and Tables 5 and 6). Table 9 shows a further analysis of the TSH receptor stimulating effects of hMAb TSHR1 IgG and Fab and the lymphocyte donor serum IgG.

The effects of porcine TSH and hMAb TSHR1 IgG on stimulation of cyclic AMP production in CHO cells expressing the TSH receptor were additive as can be seen in the results shown in Table 10.

Typical results observed in the stimulation of cyclic AMP assay with the reference preparation NIBSC 90/672 are shown in Table 11.

Overall our analysis indicates that the human monoclonal autoantibody hMAb TSHR1 has the TSH receptor binding and thyroid stimulating characteristics of the TSH receptor autoantibodies in the serum of the lymphocyte donor.

### **CONCLUSIONS**

- (a) We have produced a human monoclonal autoantibody to the TSH receptor which has similar properties to the TSH receptor autoantibody in the donor patient's serum.
- (b) The monoclonal antibody IgG and Fab preparations are powerful thyroid stimulators and effective inhibitors of labelled TSH binding to the TSH receptor.
- (c) Binding of labelled MAb IgG and Fab preparations to the TSH receptor is inhibited by TSH receptor autoantibody positive sera from patients with Graves' disease but not by healthy blood donor sera or sera from patients with other autoimmune diseases.
- (d) TSH receptor autoantibodies which act as TSH antagonists as well as TSH receptor autoantibodies which act as TSH agonists inhibit labelled hMAb TSHR1 binding to the TSH receptor.
- (e) hMAb TSHR1 preparations coated onto plastic tubes bind TSH receptor and this binding is inhibited by TSH receptor autoantibodies in different patient sera.
- (f) These results indicate that hMAb TSHR1 and/or its derivatives can be used as a replacement for TSH in
  - (i) assays for TSH receptor autoantibodies, TSH and related ligands
  - (ii) various in vivo applications involving provision of TSH agonist or TSH antagonist activities.

(iii) identification and provision of new types of TSH receptor autoantibody binding sites.

Table 1 Effect of patient sera on <sup>125</sup>I-labelled hMAb TSHR1 IgG binding to the TSH receptor and comparison with effect on <sup>125</sup>I-labelled TSH binding to the TSH receptor

Test material	inhibition of labelled hMAb	inhibition of TSH	Test material	inhibition of labelled hMAb	inhibition of TSH
	TSHR1 binding	binding	·	TSHR1 binding	binding
G1	62	80	N1	3.1	7.7
G2	91	93	N2	2.4	2.6
G3	91	76	N3	-1.0	4.5
G4	94	92	N4	-11	6.5
G5	93	94	N5	1.7	5.0
G6	76	85	N6	2.8	1.7
G7	87	90	N7	5.2	-0.8
G8	65	45	N8 .	3.5	0.2
G9	88	90	N9	2.8	-0.6
G10/10	83	59	N10	4.5	2.2
G10/20	69	43	D1	-4.8	2.2
G10/40	56	29	A1	-3.1	1.3
G10 /80	42	19	A2	-3.5	-3.0
G11/10	75	73			
G11/20	59	54			
G11 /40	39	33			
G11 /80	22	18			

G1-G11 are sera from patients with a history of Graves' disease.

G9 serum has high levels of TSH blocking (ie TSH antagonist activity).

G10 and G11 have high levels of thyroid stimulating activity.

G10 is the lymphocyte donor serum.

/10, /20 etc indicate dilution factor in a pool of healthy blood donor sera.

N1-N10 are sera from healthy blood donors.

D1 is from a patient with type 1 diabetes mellitus (positive for autoantibodies to glutamic acid decarboxylase).

A1 and A2 are from patients with Addison's disease (positive for steroid-21-hydroxylase autoantibodies).

In the presence of the pool of healthy blood donor sera about 25% of <sup>125</sup>I-labelled MAb IgG bound to the TSHR coated tubes.

Table 2 Effect of patient sera on <sup>125</sup>I-labelled hMAb TSHR1 Fab binding to the TSH receptor and comparison with effect on <sup>125</sup>I-labelled TSH binding to the TSH receptor

	•	•
Test material	inhibition of labelled Fab binding	inhibition of TSH binding
NIBSC 90/672 diluted in a pool		
of healthy blood donor serum		. 13
to 1 U/L	17	. 13
to 2 U/L	27	24
to 4 U/L	47	44
to 8 U/L	61	65
Healthy blood donor serum A	-3	<10
Healthy blood donor serum B	3	<10
Healthy blood donor serum C	4	<10
Healthy blood donor serum D	-4	<10
Healthy blood donor serum E	. 0	<10
Graves' serum F	64	78
Graves' serum G	42	54
Graves' serum H	49	69
Graves' serum I	24	36
Graves' serum J	76	88

Table 3 Binding of TSHR to plastic tubes coated with hMAb TSHR 1 Fab and inhibition of TSHR binding by sera containing TSHR autoantibodies

Test material	cpm bound <sup>1</sup>
Healthy blood donor serum A	8406
Healthy blood donor serum B	8430
TSHR autoantibody positive serum 1	1527
TSHR autoantibody positive serum 2	1131
TSHR autoantibody positive serum 3	1199

 $<sup>^1</sup>$  TSHR binding was detected using a  $^{125}$ I-labelled mouse monoclonal antibody to the TSHR C terminus; total cpm = 39,000 per tube.

Table 4 Effect of patient serum samples on binding of biotin labelled hMAb

TSHR1 and biotin labelled TSH to ELISA plate wells coated with

TSHR

	hMAb TSI	TR1 biotin	TSH	biotin
	OD <sub>450</sub>	% inhibition	OD <sub>450</sub>	% inhibition
HBD pool	1.852	0	1.778	0
HBD pool plus 1U/mL	1.46	21	1.489	16
HBD pool plus 2U/mL	1.168	37	1.304	27
HBD pool plus 4U/mL	0.792	57	0.947	47
HBD pool plus 8U/mL	0.539	71	0.492	72
HBD pool plus 40U/mL	0.118	94	0.233	87
Serum P1	1.415	24	1.397	21
Serum P2	1.264	32	1.256	29
Serum P3	0.558	70	0.408	77
Serum P4	0.763	59	0.907	49
Serum P5	1.047	43	_	
Serum P6	0.843	55	<u> </u>	
Serum P7	1.429	23	-	
HBD 1	1.745	6	1.713	4
HBD 2	1.807	2	-	
HBD 3	1.779	4	1.626	9
HBD 4	1.821	2	-	
HBD 5	1.841	1	1.660	7
HBD 6	1.762	5	1.777	0
HBD 7	1.799	3	1.767	11
HBD 8	1.783	4	1.703	4
HBD 9	1.792	3	1.669	3

HBD = healthy blood donor serum

U/mL are units of NIBSC 90/672

Serum P1-P7 are from patients with Graves' disease

Table 5 Inhibition of TSH binding by WHO reference preparation NIBSC 90/672 and by hMAb TSHR1 IgG and Fab preparations

	Samples diluted in serum <sup>1</sup>				Samples diluted in assay buffer			
Sample	% inhibition	units/L	units/mg	mean units/mg	% inhibition	units/L	units/mg	mean units/mg
NIBSC 90/672								<u> </u>
0.125 units/L					. 2			
0.25 units/L					4			
0.5 units/L	-				• 11			
1.0 units/L	15				19		<u> </u>	
2.0 units/L	28				38			
4.0 units/L	48				64			
8.0 units/L	69				83			
40.0 units/L	95				94		<u> </u>	
hMAb TSHR1 IgG					•			
0 ng/mL	1				0			
0.3 ng/mL	1				2		<b></b>	
1 ng/mL	3				3			ļ
3 ng/mL	7				10	0.46		
10 ng/mL	21	1.48	148		33	1.73	173	
30 ng/mL	46	3.9	130	138	70	4.8	160	163
100 ng/mL	81	13.5	135		92	15.6	156	
300 ng/mL	92 .				95	>40	<del> </del>	
hMAb TSHR1 Fab								
0.3 ng/mL	5				-2			_
1 ng/mL	5				1			<u> </u>
3 ng/mL	16	1.05	351		16	0.8	265	
10 ng/mL	36	2.77	277		52	2.9	291	309
30 ng/mL	69	8.0	267	288	86	9.6	372	
100 ng/mL	89	23.7	237		92	16.9		
300 ng/mL	93				94			-
2G4 IgG <sup>2</sup>								
0.3 ng/mL	2				-3			
3 ng/mL	1				-6			
30 ng/mL	0	1			-5			
300 ng/mL	3				4			_
2G4 Fab <sup>2</sup>								
0.3 ng/mL	4				-5			
3 ng/mL	4				-6			
30 ng/mL	1 1	1			-5			
300 ng/mL	2	<del>-  </del>			-6			

<sup>1</sup> Pool of healthy blood donor serum, 14.9% of total cpm bound to the TSHR in the presence of this serum pool only. 14.7% of total cpm bound to the TSHR in the presence of buffer only.

<sup>2</sup> 2G4 is a human monoclonal autoantibody to thyroid peroxidase.

Table 6 serum IgG

Inhibition of TSH binding by lymphocyte donor serum and donor

	Sa	mples dilu	ited in serw	m <sup>1</sup>	Sam	ples dilute	ed in assay b	uffer
ımple	% inhibition	units/L²	units/mg or (units/mL in undiluted serum)	mean units/mg or (units/mL)	% inhibition	units/L²	units/mg or (units/mL in undiluted serum)	mean units/mg or (units/m L)
onor serum								
luted 1000x	6				10			
luted 300x	18	1.2	(0.36)		28	1.3	(0.39)	
luted 100x	42	3.2	(0.32)	(0.36)	62	3.9	(0.39)	(0.40)
luted 30x	78	11.3	(0.39)		91	13.5	(0.41)	
luted 10x	93	34			95	>40		
onor serum IgG								
mg/mL	0	0			0			
01 mg/mL	7 .				19	0.87		
03 mg/mL	23	1.6	0.053		37	1.9	0.063	
1 mg/mL	57	5.1	0.051	0.054	78	6.4	0.064	0.063
3 mg/mL	85	17	0.057		93	19	0.063	
mg/mL	96	43			96	>40		
ealthy blood								
luted 1000x	0				3			
luted 1000x	1	<del> </del>	<del> </del>	<del> </del>	4			<del> </del>
luted 100x	1	<del></del>			11			
lutou Tox								
ealthy blood								
onor pool serum					1			
<u>:</u> G								<u> </u>
01 mg/mL	2	1			2	ļ		<b>_</b>
1 mg/mL	1		•		5		<u> </u>	ļ
mg/mL	3	<u> </u>			7		1	

<sup>&</sup>lt;sup>1</sup> Pool of healthy blood donor serum, 14.7% of total cpm bound to the TSHR in the presence of this serum pool only. 16.3% of total cpm bound to the TSHR in the presence of buffer only.

<sup>&</sup>lt;sup>2</sup> Units shown are NIBSC 90/672 international TSHR autoantibody reference preparation.

Specific activities of hMAb TSHR1 and lymphocyte donor serum and IgG preparations Table 7

	Inhibition of TS	SH binding assay	Stimulation of cyclic AMP assa		
Preparation	Units/mg <sup>1,2</sup>	Units/nmole <sup>1,2</sup>	Units/mg <sup>1</sup>	Units/nmole <sup>1</sup>	
hMAb TSHR1 IgG	150	22	180	26	
	300	15	700	35	
hMAb TSHR1 Fab		0.000	0.33	0.048	
Donor serum IgG	0.059	0.009	0.33	1.0	
Donor serum units/mL	0.38			1.8	

<sup>&</sup>lt;sup>1</sup> Units shown are NIBSC 90/672.
<sup>2</sup> Values are a mean of results obtained in serum and in assay buffer (see Tables 4 and

Table 8 Summary of specific activities of hMAb TSHR1 and lymphocyte donor serum and serum IgG determined in several stimulation of cyclic AMP assays

Preparation	Mean Units per mg	Number of determinations	Standard Deviation
hMAb TSHR1 IgG	318	16	189
hMAb TSHR1 Fab	492	10	184
Donor serum IgG	0.10	10	0.08
Donor serum	0.9	4	0.6

Table 9 Further analysis of the effects of hMAb TSHR1 IgG and Fab and lymphocyte donor serum IgG in the cyclic AMP stimulation assay

Sample	Mean cyclic AMP	number of determinations	Standard Deviation
	per well (pmols)	determinations	Deviation
1MAb TSHR1 IgG	. •		
0ng/mL	0.96	6	0.048
0.3ng/mL	1.25	. 6	0.12
1ng/mL	1.84	6	0.16
3ng/mL	3.4	5	0.37
10ng/mL	6.6	5	0.62
hMAb TSHR1 Fab			
0ng/mL	0.60	6	0.068
0.3ng/mL	1.11	6	0.11
1ng/mL	1.99	6	0.39
3ng/mL	4.9	6	0.44
10ng/mL	10.6	6	0.86
Control human MAb (2G4) <sup>1</sup>			
IgG 0ng/mL	0.72	11	0.19
IgG 10ng/mL	0.61	11	0.16
Fab 10ng/mL	0.61	4	0.044
Lymphocyte donor serum IgG			
3μg/mL	1.67	6	0.38
10μg/mL	4.20	6	0.93
30μg/mL	6.22	6	0.73
Healthy blood donor pool serum Igo	<u>G</u>		

30μg/mL	0.38	6	0.10
1			

<sup>&</sup>lt;sup>1</sup> 2G4 is a human monoclonal autoantibody to thyroid peroxidase

Table 10 Additive effect of TSH and hMAb TSHR1 IgG in stimulation of cyclic

AMP assays

	Experiment 1			Experiment 2			
	Sample	cyclic AMP <sup>1</sup> (pmols per well)		Sample	cyclic AMP <sup>1</sup> (pmols per well)		
A	Buffer only	0.57	A	Buffer only	0.42		
В	Porcine TSH 0.1ng/mL	1.07	В	Porcine TSH 0.05ng/mL	1.07		
·C	hMAb TSHR1 1ng/mL	1.41	C	hMAb TSHR1 0.5ng/mL	0.92		
Bp	olus C	2.08	Вр	lus C	1.92		

<sup>&</sup>lt;sup>1</sup> Values shown are means of closely agreeing duplicate determinations

Table 11 Effects of NIBSC 90/672 in the cyclic AMP stimulation assay

Sample	Mean cyclic AMP per well (pmols)	number of determinations	Standard Deviation
Buffer only	0.60	6	0.068
0.1 units/L	1.09	6	. 0.085
0.3 units/L	1.49	5	0.11
1.0 units/L	3.52	5	0.46
3.0 units/L	8.16	6	1.39

## Sequence Listings

### SEQ ID NO. 1

QMQLVQSGAEVKKPGESLKISCRGSGYRFTSYWINWVRHVPGKGLEWMGRI DPTDSYTNYSPSFKGHVTVSADKSINTAYLQWSSLKASDTGMYYCARLEPGY SSTWSVNWGQGTLVTVSS

SEQ ID NO. 2

**SYWIN** 

SEQ ID NO. 3

RIDPTDSYTNYSPSFKG

SEQ ID NO. 4

**LEPGYSSTWSVN** 

SEQ ID NO. 5

QMQLVQSGAEVKKPGESLKISCRGSGYRFTSYWINWVRHVPGKGLEWMGRI DPTDSYTNYSPSFKGHVTVSADKSINTAYLQWSSLKASDTGMYYCARLEPGY SSTWSVNWGQGTLVTVSSASTKGPSVFP

### SEQ ID NO. 6

LPVLTQPPSVSGAPRQRVTISCSGNSSNIGNNAVNWYQQLPGKAPKLLIYYDD QLPSGVSDRFSGSRSGTSASLAIRGLQSEDEADYYCTSWDDSLDSQLFGGGTR LTVLG

SEQ ID NO. 7

**SGNSSNIGNNAVN** 

SEQ ID NO. 8

YDDQLPS

SEQ ID NO. 9

TSWDDSLDSQL

**SEQ ID NO. 10** 

caaatgcagctggtgcagtctggagcagaggtgaaaaagcccggggagtctctgaagatctcctgtaggggttctggata caggtttaccagctactggatcaactgggtgcgccacgtgcccgggaaaggcctagagtggatgggcaggattgatccta ctgactcttataccaactacagtccatccttcaaaggccacgtcaccgtctcagctgacaagtccatcaacactgcctacctg cagtggagcagcctgaaggcctcggacaccggcatgtattactgtgcgaggctcgaaccgggctatagcagcacctggt ccgtaaattggggccagggaaccctggtcaccgtctcctca

**SEQ ID NO. 11** 

agctactggatcaac

SEQ ID NO. 12

aggattgatcctactgactcttataccaactacagtccatccttcaaaggc

**SEQ ID NO. 13** 

ctcgaaccgggctatagcagcacctggtccgtaaat

**SEQ ID NO. 14** 

caa at geaget get geaget et geage ag ag get gaa aa a ag ee eg gegag get et et gaag at et eet gegag get et gegag gede et gegag gede en gegag gegag gede en gegag gegag

63

ctgactcttataccaactacagtccatccttcaaaggccacgtcaccgtctcagctgacaagtccatcaacactgcctacctg cagtggagcagcctgaaggcctcggacaccggcatgtattactgtgcgaggctcgaaccgggctatagcagcacctggt ccgtaaattggggccagggaaccctggtcaccgtctcctcagcctccaccaagggcccatcggtcttcccc

### **SEQ ID NO. 15**

ctgcctgtgctgactcagccaccctcggtgtctggagccccaggcagagggtcaccatctctgttctggaaacagctcc
aacatcggaaataatgctgtaaactggtaccagcagctcccaggaaaggctcccaaactcctcatttattatgatgatcaact
gccctcaggggtctctgaccgattctctggctccaggtctggcacctccgcctccctggccatccgtgggctccagtctgag
gatgaggctgattattactgtacatcatgggatgacagcctggatagtcaactgttcggcggagggaccaggctgaccgtc
ctaggt

### **SEQ ID NO. 16**

tetggaaacagetecaacateggaaataatgetgtaaac

## **SEQ ID NO. 17**

tatgatgatcaactgccctca

### **SEQ ID NO. 18**

acatcatgggatgacagcctggatagtcaactg

#### **CLAIMS**

- 1. A binding partner for the TSH receptor, which binding partner comprises, or is derived from, a human monoclonal or recombinant antibody, or one or more fragments thereof, reactive with the TSH receptor.
- 2. A binding partner for the TSH receptor, which binding partner comprises, or is derived from, a human monoclonal antibody, or one or more fragments thereof, reactive with the TSH receptor.
- 3. A human monoclonal antibody, or one or more fragments thereof, reactive with the TSH receptor.
- 4. A binding partner according to any of claims 1 to 3, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 5. A binding partner according to claim 4, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 6. A binding partner according to any of claims 1 to 5, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 7. A binding partner according to claim 6, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 8. A binding partner according to any of claims 1 to 7, characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

- 9. A binding partner according to claim 8, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

- 10. A binding partner according to any of claims 1 to 9, which comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg.
- 11. A binding partner according to claim 10, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg.
- 12. A binding partner according to any of claims 1 to 11, which comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, characterised by a stimulatory activity with respect to

cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg.

- 13. A binding partner according to claim 12, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 400 units of International Standard NIBSC 90/672 per mg.
- 14. A binding partner according to any of claims 1 to 13, which comprises or is derived from one or more fragments of a monoclonal or recombinant antibody reactive with the TSH receptor, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg.
- 15. A binding partner according to claim 14, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 400 units of International Standard NIBSC 90/672 per mg.
- 16. A binding partner for the TSH receptor which comprises an antibody  $V_H$  domain selected from the group consisting of a  $V_H$  domain as shown in SEQ ID NO. 1 and a  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4.
- 17. A binding partner for the TSH receptor which comprises an antibody  $V_H$  domain as shown in SEQ ID NO. 1.

- 18. A binding partner for the TSH receptor which comprises an antibody  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4.
- 19. A binding partner for the TSH receptor which comprises:

an antibody V<sub>H</sub> domain selected from the group consisting of:

a  $V_H$  domain as shown in SEQ ID NO. 1 and a  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4; and / or

an antibody V<sub>L</sub> domain selected from the group consisting of:

a  $V_L$  domain as shown in SEQ ID NO. 6 and a  $V_L$  domain comprising one or more  $V_L$  CDRs with an amino acid sequence selected from SEQ ID NO. 7, SEQ ID NO. 8 and SEQ ID NO. 9.

- 20. A binding partner according to claim 19, comprising an antibody  $V_H$  domain as shown in SEQ ID NO. 1 paired with an antibody  $V_L$  domain as shown in SEQ ID NO. 6 to provide an antibody binding site, comprising both said  $V_H$  and  $V_L$  domains for the TSH receptor.
- 21. A binding partner according to claim 19, which comprises:

an antibody V<sub>H</sub> domain comprising:

a  $V_H$  domain comprising one or more  $V_H$  CDRs with an amino acid sequence selected from SEQ ID NO. 2, SEQ ID NO. 3 and SEQ ID NO. 4; and / or

an antibody  $V_L$  domain comprising:

- a  $V_L$  domain comprising one or more  $V_L$  CDRs with an amino acid sequence selected from SEQ ID NO. 7, SEQ ID NO. 8 and SEQ ID NO. 9.
- 22. A further binding partner capable of binding to the TSH receptor and which competes for binding to the TSH receptor with a binding partner for the TSH receptor according to any of claims 1 to 21, which further binding partner does not comprise TSH or a human antibody to the TSH receptor.
- 23. A further binding partner according to claim 22, which comprises a further antibody having a binding site for an epitope region of the TSH receptor and which competes for binding to the TSH receptor with a binding partner according to any of claims 1 to 21.
- 24. A further binding partner capable of binding to the TSH receptor and which competes for binding to the TSH receptor with a binding partner for the TSH receptor according to any of claims 16 to 21, which further binding partner comprises, or is derived from, a human monoclonal or recombinant antibody, or one or more fragments thereof, reactive with the TSH receptor.
- 25. A further binding partner according to claim 24, which comprises a further antibody having a binding site for an epitope region of the TSH receptor and which competes for binding to the TSH receptor with a binding partner according to any of claims 16 to 21.
- 26. A further binding partner according to any of claims 22 to 25, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 27. A further binding partner according to claim 26, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.

- 28. A further binding partner according to any of claims 22 to 27, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 29. A further binding partner according to claim 28, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 30. A further binding partner according to any of claims 22 to 29, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

- 31. A further binding partner according to claim 30, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

32. A polynucleotide comprising:

- (i) a nucleotide sequence as shown in SEQ ID NO. 10, SEQ ID NO. 11, SEQ ID NO. 12, SEQ ID NO. 13, SEQ ID NO. 15, SEQ ID NO. 16, SEQ ID NO. 17 or SEQ ID NO. 18, encoding an amino acid sequence of an antibody V<sub>H</sub> domain, V<sub>L</sub> domain, or CDR, as shown in SEQ ID NO. 1, SEQ ID NO. 2, SEQ ID NO. 3, SEQ ID NO. 4, SEQ ID NO. 6, SEQ ID NO. 7, SEQ ID NO. 8 or SEQ ID NO. 9;
- (ii) a nucleotide sequence encoding a binding partner for the TSH receptor according to any of claims 16 to 21, or encoding an amino acid sequence of an antibody V<sub>H</sub> domain, V<sub>L</sub> domain, or CDR, of a binding partner for the TSH receptor according to any of claims 16 to 21;
- (iii) a nucleotide sequence differing from any sequence of (i) in codon sequence due to the degeneracy of the genetic code;
- (iv) a nucleotide sequence comprising an allelic variation of any sequence of(i);
- (v) a nucleotide sequence comprising a fragment of any of the sequences of (i), (ii), (iii), or (iv) and in particular a nucleotide sequence comprising a fragment of any of the sequences of (i), (ii), (iii), (iv) or (v) and encoding a Fab fragment, a Fd fragment, a Fv fragment, a dAb fragment, an isolated CDR region, F(ab')2 fragments or a scFv fragment, of a binding partner for the TSH receptor according to any of claims 16 to 21;
- (vi) a nucleotide sequence differing from the any sequence of (i) due to mutation, deletion or substitution of a nucleotide base and encoding a binding partner for the TSH receptor according to any of claims 16 to 21, or encoding an amino acid sequence of an antibody V<sub>H</sub> domain, V<sub>L</sub> domain, or CDR, of a binding partner for the TSH receptor according to any of claims 16 to 21.

- 33. A biologically functional vector system which carries a polynucleotide according to claim 32 and which is capable of introducing the polynucleotide into the genome of a host organism.
- 34. A host cell which is transformed with a polynucleotide according to claim 32.
- 35. A process of providing a human monoclonal antibody to the TSH receptor, which process comprises:
  - (i) providing a source of lymphocytes from a subject, which subject has TSH receptor antibody activity of greater than about 0.04 units of NIBSC 90/672 per mL of serum with respect to inhibition of TSH binding to the TSH receptor;
  - (ii) isolating lymphocytes from said lymphocyte source of (i);
  - (iii) immortalising the isolated lymphocytes; and
  - (iv) cloning the immortalised lymphocytes so as to produce an immortalised colony secreting a human monoclonal antibody to the TSH receptor.
  - 36. A process of providing a human monoclonal antibody to the TSH receptor, which comprises:
    - (i) providing a source of lymphocytes from a subject, which subject has TSH receptor antibody activity of greater than about 0.1 units of NIBSC 90/672 per mL of serum with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor;
    - (ii) isolating lymphocytes from said lymphocyte source of (i);
    - (iii) immortalising the isolated lymphocytes; and

- (iv) cloning the immortalised lymphocytes so as to produce an immortalised colony secreting a human monoclonal antibody to the TSH receptor.
- 37. A process according to claim 35 or 36, which comprises isolating lymphocytes from peripheral blood, thyroid tissue, spleen tissue, lymph nodes or bone marrow.
- 38. A process according to claim 35, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels of greater than about 0.1 units of NIBSC 90/672 per mL of serum with respect to inhibition of TSH binding to the TSH receptor.
- 39. A process according to claim 38, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels of greater than about 0.2 units of NIBSC 90/672 per mL of serum with respect to inhibition of TSH binding to the TSH receptor.
- 40. A process according to claim 39, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels in the range of about 0.3 to 0.5 units of NIBSC 90/672 per mL of serum with respect to inhibition of TSH binding to the TSH receptor.
- 41. A process according to claim 36, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels of greater than about 0.3 units of NIBSC 90/672 per mL of serum with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor.
- 42. A process according to claim 41, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels of greater than about 0.5 units of NIBSC 90/672 per mL of serum with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor.
- 43. A process according to claim 42, wherein the source of lymphocytes is characterised as being obtained from a subject having TSH receptor antibody levels in

the range of about 0.5 to 1.0 units of NIBSC 90/672 per mL of serum with respect to stimulatory activity of cAMP production by cells expressing the TSH receptor.

- 44. A process according to any of claims 35 to 43, which comprises infecting the isolated lymphocytes with Epstein Barr virus, and the thus immortalised lymphocytes are fused with a mouse or human cell line.
- 45. A human monoclonal antibody to the TSH receptor obtained by a process according to any of claims 35 to 44.
- 46. A human monoclonal antibody according to claim 45, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 47. A human monoclonal antibody according to claim 46, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 48. A human monoclonal antibody according to any of claims 45 to 47, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 49. A human monoclonal antibody according to claim 48, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg, or one or more fragments thereof.
- 50. A human monoclonal antibody according to any of claims 45 to 49, characterised by:

- (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 15 units of International Standard NIBSC 90/672 per mg; and
- (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

- 51. A human monoclonal antibody according to claim 50, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 120 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg;

or one or more fragments thereof.

- 52. One or more fragments of a human monoclonal antibody according to any of claims 45 to 51, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg.
- 53. One or more fragments according to claim 52, characterised by an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg.
- 54. One or more fragments of a human monoclonal antibody according to any of claims 45 to 51, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg.

- 55. One or more fragments according to claim 54, characterised by a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 400 units of International Standard NIBSC 90/672 per mg.
- 56. One or more fragments according to any of claims 52 to 55, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 30 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 50 units of International Standard NIBSC 90/672 per mg.
- 57. One or more fragments according to claim 56, characterised by:
  - (i) an inhibitory activity with respect to TSH binding to the TSH receptor, of at least about 240 units of International Standard NIBSC 90/672 per mg; and
  - (ii) a stimulatory activity with respect to cAMP production by cells expressing the TSH receptor, of at least about 400 units of International Standard NIBSC 90/672 per mg.
- 58. A process according to any of claims 35 to 44, which further comprises a further process stage whereby the obtained human monoclonal antibody is subjected to further processing techniques so as to obtain a further binding partner according to any of claims 24 to 31.
- 59. A further binding partner according to any of claims 24 to 31, obtained by a process according to claim 58.
- 60. A method of screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to the TSH receptor, said method comprising:

- (a) providing said sample of body fluid from said subject;
- (b) providing one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 and a second molecule of said binding pair comprises a binding region with which said binding partner or further binding partner interacts;
- (c) contacting said sample with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) autoantibodies to the TSH receptor present in said sample, or (ii) said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31; and
- (d) monitoring the interaction of said second molecule of said binding pair with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.
- 61. A method of screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to the TSH receptor, said method comprising:
  - (a) providing said sample of body fluid from said subject;
  - (b) contacting said sample with (i) a full length TSH receptor, or one or more epitopes thereof or a polypeptide comprising one or more epitopes of a TSH receptor, and (ii) a binding partner for the TSH receptor or further binding partner for the TSH receptor according to any of claims 1 to 31, under conditions that allow interaction of the TSH receptor with autoantibodies produced in response to the TSH receptor, so as to permit said TSH receptor, or said one or more epitopes thereof or said polypeptide, to interact with either

autoantibodies to the TSH receptor present in said sample, or said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31; and

- (c) monitoring the interaction of said TSH receptor, or said one or more epitopes thereof or said polypeptide, with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.
- 62. A method according to claim 60 or 61, which further employs one or more competitors that compete in the interaction of the binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 and the second molecule of the binding pair of claim 60 or said TSH receptor, or said one or more epitopes thereof or said polypeptide of claim 61.
- 63. A kit for screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to the TSH receptor, said kit comprising:
  - (a) one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 and a second molecule of said binding pair comprises a binding region with which said binding partner or further binding partner interacts;
  - (b) means for contacting said sample of body fluid from said subject with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) autoantibodies to the TSH receptor present in said sample, or (ii) said binding partner for the TSH receptor or further binding partner for the TSH receptor according to any of claims 1 to 31; and

- (c) means for monitoring the interaction of said second molecule of said binding pair with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.
- 64. A kit for screening for autoantibodies to the TSH receptor in a sample of body fluid obtained from a subject suspected of suffering from, susceptible to, having or recovering from autoimmune disease associated with an immune reaction to the TSH receptor, said kit comprising:
  - (a) a full length TSH receptor, or one or more epitopes thereof or a polypeptide comprising one or more epitopes of the TSH receptor;
  - (b) a binding partner for the TSH receptor or further binding partner for the TSH receptor according to any of claims 1 to 31;
  - (c) means for contacting said sample of body fluid from said subject, said TSH receptor, or said one or more epitopes thereof or said polypeptide, and said binding partner for the TSH receptor or further binding partner for the TSH receptor according to any of claims 1 to 31, under conditions that allow interaction of the TSH receptor with autoantibodies produced in response to the TSH receptor, so as to permit said TSH receptor, or said one or more epitopes thereof or said polypeptide, to interact with either autoantibodies to a TSH receptor present in said sample, or said binding partner for the TSH receptor or further binding partner for the TSH receptor according to any of claims 1 to 31; and
  - (d) means for monitoring the interaction of said TSH receptor, or said one or more epitopes thereof or said polypeptide, with said autoantibodies present in said sample, thereby providing an indication of the presence of said autoantibodies to the TSH receptor in said sample.
- 65. A kit according to claim 63 or 64, which further comprises one or more competitors that compete in the interaction of a binding partner or further binding

partner for the TSH receptor according to any of claims 1 to 31 and the second molecule of the binding pair of claim 63 or said TSH receptor, or said one or more epitopes thereof or said polypeptide of claim 64.

- 66. A method of assaying TSH and related ligands, said method comprising:
  - (a) providing a sample suspected of containing or containing TSH or related ligands;
  - (b) providing one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;
  - (c) contacting said sample with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) TSH or related ligands present in said sample, or (ii) said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31; and
  - (d) monitoring the interaction of said second molecule of said binding pair with TSH or related ligands present in said sample, thereby providing an indication of the presence of TSH or related ligands in said sample.
  - 67. A kit for assaying TSH or related ligands, said kit comprising:
    - (a) one or more pairs of binding molecules, wherein a first molecule of said binding pair comprises a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 and a second molecule of said binding pair comprises a binding region with which said binding partner interacts;

- (b) means for contacting a sample suspected of containing or containing TSH or related ligands with said one or more pairs of binding molecules so as to permit said second molecule of said binding pair to interact with either (i) TSH or related ligands present in said sample, or (ii) said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31; and
- (c) means for monitoring the interaction of said second molecule of said binding pair with TSH or related ligands present in said sample, thereby providing an indication of the presence of TSH or related ligands in said sample.
- A process of identifying one or more epitope regions of the TSH receptor, which process comprises contacting a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 with a full length TSH receptor, or one or more fragments thereof, so as to allow interaction of said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 with said full length TSH receptor, or said one or more fragments thereof, and identifying the amino acids of said full length TSH receptor, or said one or more fragments thereof, with which said binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 interacts.
- 69. One or more anti-idiotypic antibodies generated to a binding region of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31.
- 70. A method of identifying antibody binding sites, which method comprises screening of phage-displayed random libraries with a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31.
- 71. A binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 for use in therapy.

- 72. A method of treating autoimmune disease associated with an immune reaction to the TSH receptor in a subject, comprising administering to said subject a therapeutically effective amount of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31.
- 73. A pharmaceutical composition comprising a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, together with one or more pharmaceutically acceptable carriers, diluents or excipients therefor.
- 74. A method of treating autoimmune disease associated with an immune reaction to the TSH receptor in a subject, comprising administering to said subject a therapeutically effective amount of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, which binding partner or further binding partner for the TSH receptor stimulates the TSH receptor.
- 75. Use of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, in the manufacture of a medicament for use in stimulating thyroid tissue, or tissue containing a TSH receptor.
- 76. Use of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, in the manufacture of a medicament for use in the treatment of thyroid cancer.
- 77. A method of stimulating thyroid tissue, and / or tissue containing the TSH receptor, which method comprises administering to a patient in need of such stimulation a diagnostically or therapeutically effective amount of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31.
- 78. In combination a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, together with one or more further agents capable of stimulating thyroid tissue, and / or tissue containing a TSH receptor, for simultaneous, separate or sequential use in stimulating thyroid tissue, and / or tissue containing a TSH receptor.

- 79. A combination according to claim 78, wherein said one or more further agents comprise recombinant human TSH and / or one or more variants, analogs, derivatives or fragments thereof, or variants, analogs or derivatives of such fragments.
- 80. A combination according to claim 79, wherein said one or more further agents acts independently of binding to the TSH receptor.
- 81. A method of treating autoimmune disease associated with an immune reaction to a TSH receptor in a subject, comprising administering to said subject a therapeutically effective amount of a binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, which binding partner or further binding partner for the TSH receptor inactivates or renders unresponsive the TSH receptor to TSH, TSH receptor autoantibodies or other stimulators.
- 82. A binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31 for use as a replacement source for patient serum required to contain TSH receptor antibody or antibodies.
- 83. A binding partner or further binding partner for the TSH receptor according to any of claims 1 to 31, for use in a preparation required to comprise a defined concentration of TSH receptor antibody or antibodies.

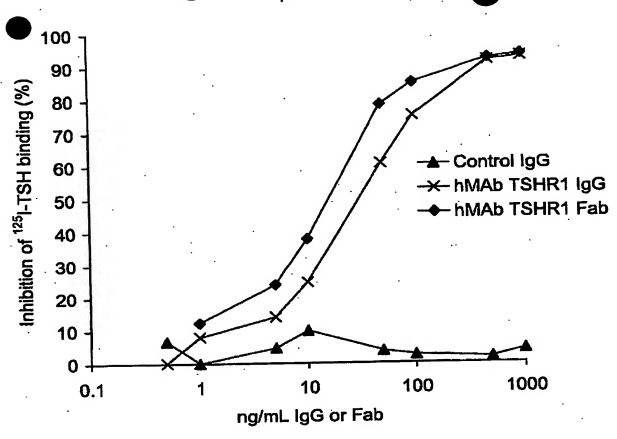
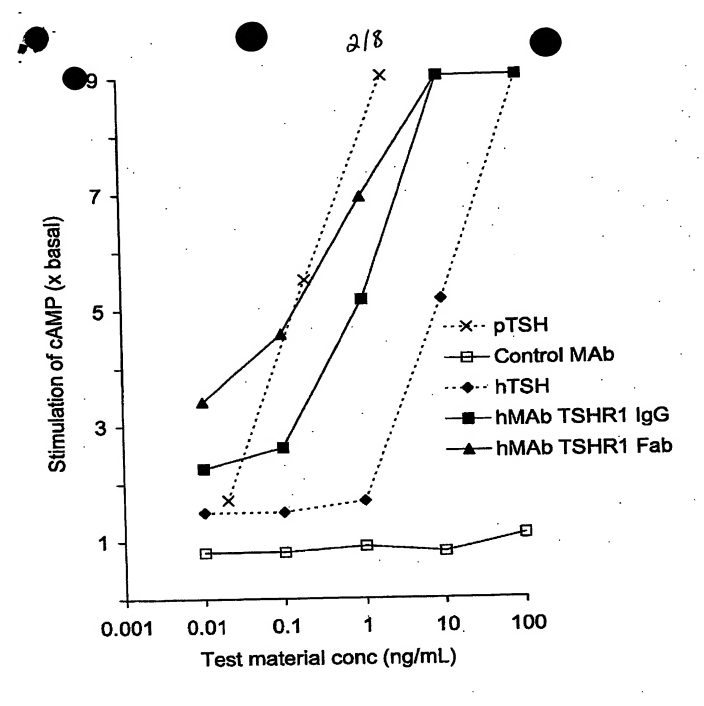


Figure 1 Inhibition of labelled TSH binding to TSHR coated tubes by hMAb

TSHR1 IgG and Fab. The control IgG was a human monoclonal autoantibody to GAD<sub>65</sub>.



Thyroid stimulating activities of hMAb TSHR1 IgG and Fab, porcine TSH (70 units/mg; pTSH), recombinant human TSH (6.7 units/mg; hTSH) and a control monoclonal antibody (MAb: a human monoclonal autoantibody to thyroid peroxidase (2G4)). Basal = cAMP produced in the presence of NaCl free Hanks Buffered Salt Solution only.

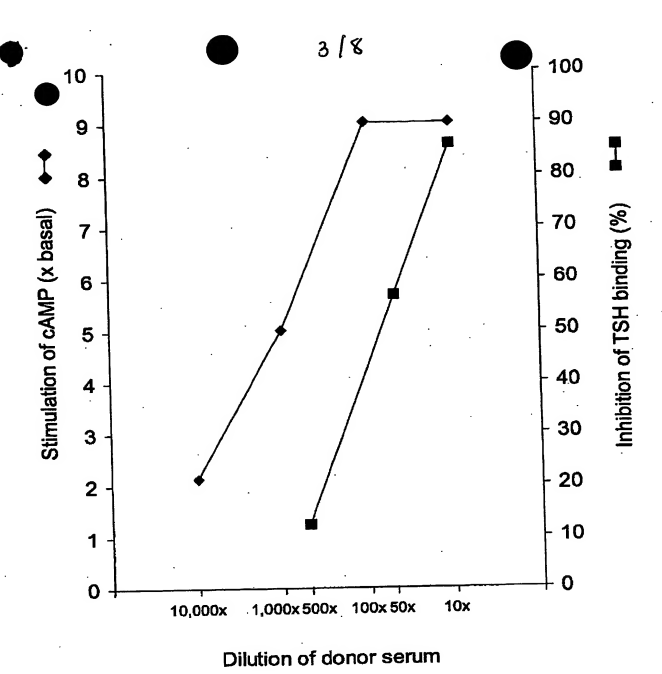
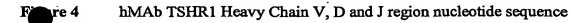


Figure 3 Effect of lymphocyte donor serum on inhibition of TSH binding to the TSHR and on stimulation of cyclic AMP in TSHR transfected CHO cells. In the case of the binding inhibition assay the serum was diluted in a pool of healthy blood donor sera. For the stimulation assay, the serum was diluted in NaCl free Hanks Buffered Salt Solution. Healthy blood donor sera (n = 3) gave responses ranging from  $1.1 - 1.3 \times 1.3 \times$ 



#### Figure 4a

caaatgcagctggtgcagtctggagcagaggtgaaaaagcccggggagtc
tctgaagatctcctgtaggggttctggatacaggtttaccagctactgga
tcaactgggtgcgccacgtgcccgggaaaggcctagagtggatgggcagg
attgatcctactgactcttataccaactacagtccatccttcaaaggcca
cgtcaccgtctcagctgacaagtccatcaacactgcctacctgcagtgga
gcagcctgaaggcctcggacaccggcatgtattactgtgcgaggctcgaa
ccgggctatagcagcacctggtccgtaaattggggccagggaaccctggt
caccgtctcctcagcctccaccaagggcccatcggtcttcccc

# Figure 4b

caaatgcagctggtgcagtctggagcagaggtgaaaaagcccggggagtc PCR primer	50
tetgaagateteetgtaggggttetggataeaggtttaee <mark>agetaetgga</mark>	100
teaactgggtgcgccacgtgcccgggaaaggcctagagtggatgggc	150
attgatectactgactettataccaactacagt@catecttcaaaggeca	200
cgtcaccgtctcagctgacaagtccatcaacactgcctacctgcagtgga	250
gcagcctgaaggcctcggacaccggcatgtattactgtgcgaggctcgaa	300
CDR III ccgggctatagcagcacctggtcgtaaattggggccagggaaccctggt	350
constant region	
caccgtctcctca <b>gcctccaccaagggcccatcggtcttccccc</b>	394

## Figure 5 hMAb TSHR1 Heavy Chain V, D and J region amino acid sequence

Figure 5a

QMQLVQSGAEVKKPGESLKISCRGSGYRFTSYWINWVRHVPGKGLEWMGR

IDPTDSYTNYSPSFKGHVTVSADKSINTAYLQWSSLKASDTGMYYCARLE

PGYSSTWS VNWGQGTLVTVSSASTKGPSVFP

Figure 5b

OMOLVOSGAEVKKPGESLKISCRGSGYRFT COR I	50
EDETDSYTMYSESEKGHVTVSADKSINTAYLQWSSLKASDTGMYYCARLE CDR II	100
PGYSSEWS VNWGQGTLVTVSS ASTKGPSVFP constant region	131

### Figure 6

### hMAb TSHR1 Light Chain DNA sequence

#### Figure 6a

ctgcctgtgctgactcagccaccctcggtgtctggagccccaggcagag
ggtcaccatctcctgttctggaaacagctccaacatcggaaataatgctg
taaactggtaccagcagctcccaggaaaggctcccaaactcctcatttat
tatgatgatcaactgccctcaggggtctctgaccgattctctggctccag
gtctggcacctccgcctccctggccatccgtgggctccagtctgaggatg
aggctgattattactgtacatcatgggatgacagcctggatagtcaactg
ttcggcggagggaccaggctgaccgtcctaggt

#### Figure 6b

•	
<pre>ctgcctgtgctgactcagccaccctcggtgtctggagcccccaggcagag PCR primer</pre>	50
ggtcaccatctcctgttctggaaacagctccaacatcggaaataatgetg	100
CDC1	
taaactggtaccagcagctcccaggaaaggctcccaaactcctcatttat	150
fatgatgatcaactgccctcaggggtctctgaccgattctctggctccag	200
CDR II	
gtctggcacctccgctccctggccatccgtgggctccagtctgaggatg	250
·	
aggetgattattaetgtacatcatgggatgacageetggatagteaaetg	300
CDR III	
	333
ttcggcggagggaccaggctgaccgtcctaggt	223

# Figure 7 hMAb TSHR1 Light Chain protein sequence

Figure 7a

LPVLTQPPSVSGAPRQRVTISCSGNSSNIGNNAVNWYQQLPGKAPKLLIY

YDDQLPSGVSDRFSGSRSGTSASLAIRGLQSEDEADYYCTSWDDSLDSQL

FGGGTRLTVLG

Figure 7b

LPVLTOPPSVSGAPRORVTISC <mark>SGNSSNIGNNÄVN</mark> WYQQLPGKAPKLLIY PCR primer CDR I	50
YDDOLPS GVSDRFSGSRSGTSASLAIRGLQSEDEADYYCESWDDSLDSQL CDR III CDR III	100
FGGGTRLTVLG	117

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